STRUCTURE, PROPERTIES AND PREPARATION OF PEROVSKITE-TYPE COMPOUNDS...

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PREFACE

Since 1945, when the ferroelectric properties of barium titanate were reported by von Hippel in the United States and independently by workers in other countries, ABO₃ compounds with the perovskite structure have been studied extensively. These studies have resulted in the discovery of many now ferroelectric and piczoelectric materials. Most of the literature written on perovskite-type compounds has been concentrated on these properties.

In addition, a number of solid-state chemists devoted many years to producing new ternary perovskite compounds of all kinds and studying their structures. By 1955 it appeared that most of the possible combinations of large A cations and smaller B ions needed to form perovskite-type compounds shad been tried. At that time, as part of a thesis problem at the University of Connecticut, I found that new perovskite-type compounds could be prepared by introducing more than one element in the B position of the perovskite structure.

At the United Aircraft Research Laboratories in 1960, J. At the United Aircraft Research Laboratories in 1960, J. (Pyle) Pinto, W. Darby and I continued this research by initiating an extensive program to study the preparation, structure and properties of these perovskite-type compounds. Because these compounds contained two different B ions with different valence states, many combinations of elements and, different valence states, many compounds were possible. These studies as well as research conducted by other workers Throughout the world resulted in large amounts of new structural and property data on perovskite-type compounds. Durtural and property data on perovskite-type compounds. Durtural and property data as a result of studies which showed compounds was generated as a result of studies which showed modulation, as thermistors, as superconductors and as inframodulation, as thermistors, as superconductors and as infra-

red windows.

The purpose of this book is to attempt to bring together the information obtained from these studies, including the

various methods of preparing powders, thin films and single crystals of perovskite-type compounds, the structure of these compounds and their properties. The properties covered are electrical conductivity, ferroelectricity, ferromagnetism, optical transmittance the electro-optical effect, catalytic properties, melting points, heats of formation, thermal expansion, densities and mechanical properties. Because of the growing number of applications for perovskite-type compounds, I felt that this information might prove valuable to applied researchers. In addition, structural data are included for scientists who are interested in correlating the structure and properties of materials.

dielectrics and Dr. Fredricks of Wright-Patterson A.F.B. for fessor R. Smoluchowski, Dr. R. Graf, Professor A. Wold and Kelly and Mrs. Nancy Letendre for their patience and effort in perovskites. I would also like to thank Dr. V. Nicolai of Charleton of Fort Monmouth, New Jersey, for discussions on ed to United Aircraft Research Laboratories, my colleagues Dr. M. Kestigian for helpful suggestions and for checking Lois, Miss Kathy Donahue, Miss Joyce Hurlburt, Mrs. Jean discussions on ordering and to Dr. John Goodenough for inand Professor P. Duwez, a member of the Advisory Committee for United Aircraft Corporation. I wish to thank Prothrough the manuscript. Finally, I am grateful to my wife, I am grateful to Professor Roland Ward and Professor Lewis Katz for introducing me to this field of research and to my previous fellow workers at the University of Connecticut for their studies on many unusual perovskite-type compounds. I must also acknowledge Professor Aaron Wold and Professor Rustum Roy for their discussions on perovskite compound preparations, Dr. Michael Kestigian and Professor A. Ŝmakula for helpful advice on crystal growing, Dr. Alexander Wells and Professor Martin Buerger for pointing out the need for a compilation and discussion of structural data of the type presented herein, Dr. Fredrick Seitz for helpful formation on the ferromagnetic properties and conductivity O.N.R. Washington, D.C., for information on lasers, Dr. reports on microwave properties of perovskites. I am indebt-J. Pinto and W. Darby; R. Fanti, Chief of Materials Sciences, n preparing this manuscript.

CHAPTER 1

INTRODUCTION

ships. An objective of this book is to point out some of these structure-property relationships as well as to provide the properties. One of the best ways of accomplishing this reader with enough data so that he can deduce some of his is to use the insight gained from structure-property relationpreparation of perovskite-type compounds. Because of the terest in obtaining these materials as optical quality single crystals. In addition, the better known ferroelectric and piezoelectric properties of perovskites have induced researchers to continue the effort to prepare them as larger and more perfect single crystals, polycrystalline compacts and thin films. Materials scientists also are continuously trying to prepare new perovskite compounds with new and improved THIS book contains details on the structure, properties, and growing number of applications for these compounds, infor-K(Tao.esNbo.ss)Os, for example, have caused considerable inmation on their preparation is becoming more in demand. The long fluorescence lifetimes observed for Cr3+ in LaAlO₃, and the large room-temperature electro-optical effect in

In this book, the oxide phases have been divided into two types, the ternary ABO₃ type and their solid solutions, where A is a large metal cation and B is a smaller metal cation and the newer complex $A(B_x^*B_y^*)O_3$ type compounds where B' and B'' are two different elements in different oxidation states and x+y=1. First, the structural data are presented in a systematic manner for quick and easy reference. A chapter is included on the identification of distortions in the structure of ternary perovskite-type compounds and of ordering in the structure of complex perovskite-type compounds using X-ray diffraction techniques. The properties of the perovskite

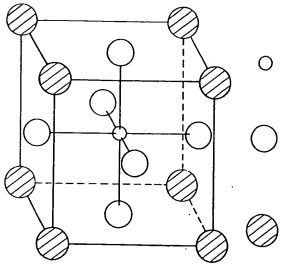
INTRODUCTION

compounds described herein are electrical conductivity, férroelectricity, ferromagnetism, optical, catalytic, melting points, heats of formation, thermal expansion and mechanical properties. Then, the preparation of these compounds as powders, thin films and single crystals are described. In addition, a chapter is included on other compounds besides oxides with the perovskite structure.

CHAPTER 2

STRUCTURE OF PEROVSKITE.TYPI COMPOUNDS

Most of the compounds with the general formula ABO₃ have the perovskite structure. The atomic arrangement in this structure was first found for the mineral perovskite, CaTiO₃. It was thought that the unit cell of CaTiO₃ could be represented by calcium ions at the corners of a cubo with titanium ions at the body center and oxygen ions at the center of the faces (Fig. 2.1). This simple cubic structure has retained the name perovskite, even though CaTiO₃ was later determined to be orthorhombic by Megaw. (1) Through the years it has been found that very few perovskite-type oxides



A CATION OXYGEN B CATION FIG. 2.1. Perovskite structure, ABOs.

have the simple cubic structure at room temperature, but many assume this ideal structure at higher temperatures.

In the perovskite structure, the A cation is coordinated with twelve oxygen ions and the B cation with six. Thus, the A cation is normally found to be somewhat larger than the B cation. In order to have contact between the A, B, and O ions, $R_A + R_O$ should equal $\sqrt{2(R_B + R_O)}$, where R_A , R_B and R_O are the ionic radii. Goldschmidt⁽²⁾ has shown that the cubic perovskite structure is stable only if a tolerance factor, t defined by $R_A + R_O = t\sqrt{2(R_B + R_O)}$, has an approximate range of 0.8 < t < 0.9, and a somewhat larger range for distorted perovskite structures. It should be noted that conflicting reports in the literature make it difficult to assign the correct unit cell dimensions for these distorted perovskite structures.

The ternary perovskite-type oxides described in this chapter will be divided into $A^{1+}B^{5+}O_{3}$, $A^{2+}B^{4+}O_{3}$, $A^{3+}B^{3+}O_{3}$ types and oxygen- and cation-deficient phases. The oxygenand cation-deficient phases will be regarded as those which contain considerable vacancies and not those phases which are only slightly non-stoichiometric. Many of these contain B ions of one element in two valence states and should not be confused with the complex perovskite compounds which contain different elements in different valence states.

The complex perovskite type compounds, $A(B_x'B_y')O_3$, will be divided into compounds which contain twice as much lower valence state element as higher valence state element, $A(B_{0.67}'B_{0.33}')O_3$, those which contain twice as much of the higher valence state element as the lower valence state element, $A(B_{0.33}'B_{0.67}')O_3$, those which contain the two B elements in equal amounts, $A(B_{0.5}'B_{0.5}')O_3$, and oxygen-deficient phases $A(B_x'B_y')O_{3-z}$.

2.1. TERNARY OXIDES

Oxides of the A1+B5+O3 Type

The A¹⁺B⁵⁺O₃ type oxides are of particular interest because of their ferroelectric properties. Potassium niobate, KNbO₃ has a structure which can be described by an ortho-

STRUCTURE OF PEROVSKITE-TYPE COMPOUND

rhombic unit cell of a=3.9714 Å, b=5.6946 Å and c=5.7203 Å where b and c equal approximately $\sqrt{2}$ a or the length of face diagonals of the simple perovskite cell, and exhibits ferroelectric properties. The sodium niobate, NaNbO₃, structure also can be described by an orthorhombic unit cell but is antiferroelectric. The unit cell is pseudotetragonal at 420° C, tetragonal at 560° C and cubic at 640° C. Unlike the structure of niobates the KTaO₃ structure is described by a cubic unit cell. The structure of NaTaO₃ is orthorhombic with the space group Pc2₁ n and all the atoms are placed in the unit cell in positions:⁽³⁾

(4a)
$$xyz$$
; \bar{x} , $y + \frac{1}{2}$, \bar{z} ; $x + \frac{1}{2}$, $y + \frac{1}{2}$, $\frac{1}{2} - z$; $\frac{1}{2} - x$, y , $z + \frac{1}{2}$

$$\begin{array}{ccccc}
x & y & z \\
y & z & y & z \\
Na & -0.01 & 0.78 & 0.02 \\
Ta & 0.50 & 0.00 & 0.00 \\
0(1) & -0.02 & 0.76 & 0.52 \\
0(2) & 0.29 & -0.03 & 0.29 \\
0(3) & 0.29 & 0.56 & 0.20
\end{array}$$

Smith and Welch⁽⁴⁾ found that potassium iodate, KIO₃, and thallous iodate, TIIO₃, also adopt the perovskite structure. Single-crystal studies showed that KIO₃ had a rhombohedral structure with unit cell parameters $\alpha = 4.410$ Å, $\alpha = 89.41$ °. Powder diffraction studies on TIIO₃ powders indicated that it also had a rhombohedral structure with cell dimensions, $\alpha = 4.510$ Å, $\alpha = 89.34$ °, while the structures of CsIO₃ and RbIO₃, on the other hand, have been reported to

Oxides of the A²⁺B⁴⁺O₃ Type

Probably the largest number of perovskite-type compounds are described by the general formula $A^{2}+B^{4}+O_{3}$, where the A cations are alkaline earth ions, cadmium or lead and the B⁴⁺ ions include Ce, Fe, Pr, Pu, Sn, Th, Hf, Ti, Zr, Mo and U. The best known compounds of this type are the titanates because of the ferroelectric properties that the barium and lead compounds exhibit. Calcium titanate, as previously mentioned, was the original example of a compound with an "ideal" cubic perovskite structure, but it was later

of strontium titanate, however, is truly cubic; the space group is Pm3m, and its atoms are in the following positions determined to have an orthorhombic structure. The structure in the unit cell

Sr:000

 $0: \frac{1}{2} \frac{1}{2} 0; \frac{1}{2} 0; \frac{1}{2} 0; \frac{1}{2}; 0; \frac{1}{2} \frac{1}{2}$

the displacements are greater in lead titanate than they are in barium titanate. For barium titanate the atoms in the because the atomic displacements in their structures produce ferroelectric properties. Neutron diffraction studies show that Barium titanate and lead titanate are of more interest unit cell are in the following positions:

Ba atom at 000

(ref. 3) Ti atom at $\frac{1}{2}$, $\frac{1}{2}$, 0.512

one O atom at $\frac{1}{2}$, $\frac{1}{2}$, 0.023

and two O atoms at $\frac{1}{2}$, 0, 0.486; 0, $\frac{1}{2}$, 0.486.

The room temperature tetragonal form of lead titanate has its atoms in these positions in its unit cell:

Pb atom at 000

(ref. 3) Ti atom at $\frac{1}{2}$, $\frac{1}{2}$, 0.541

one O atom at $\frac{1}{2}$, $\frac{1}{2}$, 0.112

and two O atoms at $\frac{1}{2}$, 0, 0.612; 0, $\frac{1}{2}$, 0.612.

While the structure of calcium titanate exhibits orthorhombic symmetry at room temperature, it becomes cubic above 900°C. Barium titanate undergoes the following transformations:

Rhombohedral —100°C orthorhombic —°C— tetragonal

cubic and lead titanate transforms

tetragonal 490°C cubic.

Roth⁽⁵⁾ regards BaZrO₃ as another compound with an "ideal" cubic perovskite structure. While his conclusion has

although Smith and Welcht felt that the SrZrO3 powder pattern should be indexed on a "doubled" cubic perovskite which has a tolerance factor of 0.86. Strontium zirconate and calcium zirconate probably have an orthorhombic structure, ance factor of 0.88 should adopt the same structure as ${
m SrTiO_3}$ been questioned, it is logical that this compound with a toler-

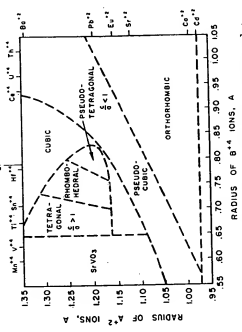


Fig. 2.2. Classification of perovskite-type compounds, A*+B*+O, (after Roth(*)).

The compound BaSnO₃ also has been reported by Smith cell. The lead zirconate structure originally was thought to be tetragonal, but was found to be orthorhombic by X-ray and neutron diffraction studies.

This selection of the unit cell was confirmed by Roth⁽⁵⁾ and Welcha to have the "ideal" cubic perovskite structure. and Megaw⁽¹⁾ independently.

but because of this new information can be assumed to be incorrect. The compound CaUO, was not found to have a it has a tolerance factor of only 0.71 it still has the perovskite structure. Roth(5) points out that a minimum tolerance factor of 0.77 was previously set for $A^2+B^4+O_3$ type compounds, cubic structure but was found to adopt the CaTiO3 structure. Another interesting compound is CaUO3, because although

A diagrammatic presentation of radius data for A²⁺B⁴⁺O₃ type compounds is shown in Fig. 2.2. The regions are deter-While this diagram holds well for compounds, there are some discrepancies in the boundaries of the ferroelectric field for solid solutions. The diagram, however, is still a useful summined from experimental data for room-temperature studies. mary of structural data.

Oxides of the A³⁺B³⁺O₃ Type

The largest number of A³⁺B³⁺O₃ type compounds were ound by Geller and Wood' to have an orthorhombic structure similar to that for GdFeO3, Fig. 2.3. The space group for these compounds is Phnm and the atoms are in the following positions:

Four Gd atoms at $\pm(x, y, \frac{1}{4}; \frac{1}{2} - x, y + \frac{1}{2}, \frac{1}{4})$ x = -0.018y = 0.060.

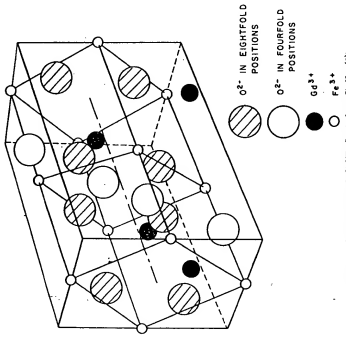


Fig. 2.3. Structure of GdFeO₃ (after Geller⁽⁶⁾).

STRUCTURE OF PEROVSKITE-TYPE COMPOUNDS

Four Fe atoms at $\frac{1}{2}$ 0 0; $\frac{1}{2}$ 0 $\frac{1}{2}$; 0 $\frac{1}{2}$ 0; 0 $\frac{1}{2}$ $\frac{1}{2}$.

Four O atoms at the same positions listed for Gd atoms, but with x = 0.05, y = 0.47

eight 0 atoms at $\pm (xyz; \frac{1}{2} - x, y + \frac{1}{2}, \frac{1}{2} - z; \hat{x}, \bar{y}, z + \frac{1}{2};$ $\frac{1}{2}-y$, \bar{z}

z = 0.05 $x = -0.29, \quad y = 0.275,$ where

the perovskite structure can be seen in Fig. 2.3. The unit cell for the GdFeO₃ structure, $a_0 = 5.346$ Å, $b_0 = 5.616$ Å, The relationship of the orthorhombic unit cell to that of c = 7.668 Å contains four distorted perovskite units.

spectively. The compounds LaAlO₃, NdAlO₃ and PrAlO₃ also have this rhombohedral structure which is probably quiteit is quite possible that many additional $A^3 + B^3 + O_3$ compounds which have been reported to have monoclinic structures pounds which were confirmed as having the GdFeO₃ strucure are EuAlO, EuFeO, GdAlO, GdCrO, GdVO, LaAlO, NdGaO3, PrCrO3, PrFeO3, PrGaO3, PrScO3, PrVO3, SmAlO3, SmCrO3, SmFeO3, YScO3, YAlO3, YCrO3, YFeO3, NdScO3 and GdScO3. Two of these compounds, LaGaO3 and SmAlO3 transform to a rhombohedral form at 900° and 850°C re-Wood(6) were able to show from single crystal studies that could really have orthorhombic structures. Some of the com-LaCrO3, LaGaO3, LaScO3, NdAlO3, NdCrO3, NdFeO3, NdVO3, Looby and Katz⁽⁷⁾ thought they had found a new type ern on the basis of a monoclinic cell, but pointed out that the correct unit cell might be orthorhombic. Geller and the structure of YCrO3 was similar to that of GdFeO3. Thus, of structure adopted by YCrO3, and indexed the powder patsimilar to that of GdFeO3.

counds according to the constituent ionic radii. All of the ype structures. Where both the A and B ions are small, the When both the A and B ions are large, the phases form ${
m La_2O_3}$ Figure 2.4 presents a classification of A^{3+B³⁺O₃ type com-} compounds in the upper left of the diagram form perovskitecompounds have the corundum- or ilmenite-type structures. ype structures.

While none of the A3+B3+O3 type compounds have the Cr3+ substitution has produced considerable interest in these compounds. Lanthanum aluminum oxide, LaAlO3, with "ideal" cubic perovskite structure, the rhombohedral perovskites such as LaAlO3 are only slightly distorted. The search for laser host materials with cubic crystallographic sites for However, the phase transition at 435° has presented consid-= $90^{\circ}4'$, has been widely studied as a laser host material. erable problems in trying to grow it in single crystal form.

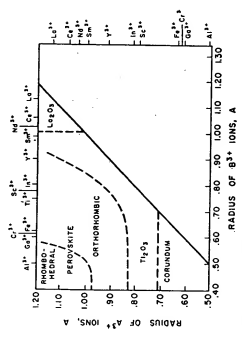


Fig. 2.4. Classification of perovskite-type compounds, $A^{3+}B^{3+}O_{8}$ (after Roth⁽⁶⁾).

Non-stoichiometric Ternary Oxides

found to have the cubic perovskite structure in the range $0.3 < x < 0.95^{(8)}$ and the phases $\text{Li}_x \text{WO}_3$ in the range $0.\overline{3}5$ Probably the best known non-stoichiometric ternary oxides are the tungsten bronzes. The phases $\mathrm{Na_{\star}WO_{3}}$ have been $< x < 0.57.^{(8)}$ The lattice constants of these materials vary er range of non-stoichiometry exists in the strontium niobium bronzes(9) where the alkaline earth metal ion mole fraction can vary from 0.7 to 0.9 and in La_xVO₃ where 0.66 < x < 1. linearly with increasing amounts of alkali metal ion. A small

Rooksby et al. (10) reported the preparation of a group of perovskite-type rare earth niobates and tantalates. The struc-

STRUCTURE OF PEROVSKITE-TYPE COMPOUNDS

ture of these $\mathrm{A}_{o.33}\mathrm{BO}_{3}$ type compounds was tetragonal, orthorhombic or monoclinic.

surprising in view of the fact that ReO₃ is stable without position without changes in structure. However, different The existence of these cation deficient compounds is not A ions. The deficiencies can be tolerated over ranges of comamounts of A ion are necessary to stabilize the structure depending on which Bion is in the octahedrally coordinated sites.

skite structure. The phases $SrBO_{3-\star}$ where B is Ti or \bar{V} have been found to have the perovskite structure over the range 0 < x < 0.5 for the titanium phases and 0 < x < 0.25 for the vanadium phases. Both $\mathrm{SrVO}_{2.75}$ and $\mathrm{SrTiO}_{2.5}$ were found to Oxygen deficiencies have also been observed in the perovhave cubic structures. Similar phases have been reported in the SrFeO3-x, (11-13) CaMnO3-x (13, 14) and SrCoO3-x (13) systems although the oxygen deficiency is not as great.

tures could be obtained in calcium perovskites, which are normally distorted, by producing oxygen vacancies. The Coates and McMillan showed that cubic perovskite strucphases CaMnO₃ and CaTiO₃ become cubic with the introduction of deficiencies. As these authors point out, studies in standing of the effect of nonstoichiometry on the perovskite this area are not abundant enough to obtain a good understructure.

2.2. COMPLEX OXIDES

Oxides of the $A^{2+}(B_{0.67}^3B_{0.33}^6)O_3$ Type

B³⁺ ions as B⁶⁺ ions is not well established. Fresia et al., (15) who prepared one of the first compounds of this type, Ba(Sco.67Wo.33)O3, felt that it probably had an ordered perovskite structure described by Steward and Rooksby. (16) In this structure the two different B ions alternate at the corners of the simple cubic unit cell of the perovskite structure so that the cell edge has to be doubled (see Fig. 2.5). The space The structure of compounds which contain twice as many group is Fm3m and the atomic positions are

B: $00\frac{1}{2}$; $\frac{1}{2}$ 00; $0\frac{1}{2}$ 0; $\frac{1}{2}$ $\frac{1}{2}$ $\frac{1}{2}$ $\frac{1}{2}$ $\frac{1}{2}$ $\frac{1}{2}$ $\frac{1}{2}$ $\frac{1}{2}$ $\frac{1}{2}$ $\frac{1}{2}$; $\frac{1}{2}$ 0 $\frac{1}{2}$ $\frac{1}{2}$; $\frac{1}{2}$ 0 $\frac{1}{2}$

O: $u \circ 0$; etc. (24 positions)

Since the B' and B'' have to be present in equal amounts in this structure, $Ba(Sc_{0.67}W_{0.33})O_3$ should probably be written

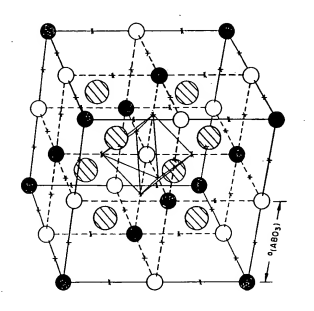




Fig. 2.5. The $(NH_4)_s$ FeF₆ structure: cubic ordered perovskite-type $A(B_{0,s}'B_{0,s}')O_s$ (after L. Pauling, J. Am. Chem. Soc. 46, 2738 (1924)).

Ba[Sc_{0.6}(Sc_{0.17}W_{0.33})]O₃ where three-quarters of the Sc ions are on the B' sites and one-quarter of the Sc ions are randomly distributed with the W atoms on the B'' sites. Sleight and Ward⁽¹⁷⁾ also found that it was necessary to use the doubled unit cell to index all observed lines in the X-ray patterns of A(B_{0.67}U_{0.33})O₃ and A(B_{0.67}Re_{0.33})O₃⁽¹⁸⁾ compounds. When

the compound Sr(Cr_{0.5}Re_{0.5})O₃ was changed in composition to obtain Sr(Cr_{0.67}Re_{0.53})O₃ no new phase appeared but the lattice expanded and the intensities of the superlattice lines in the X-ray patterns showed a marked decrease. However, no detailed structure studies have been conducted to determine the arrangement of atoms in these phases.

Oxides of the $A^{2+}(B_{0.33}^{2+}B_{0.67}^{5+})O_3$ Type

A large number of compounds containing niobium and tantalum as one of the B ions in perovskite structure and a divalent ion as the other B ion were prepared by Roy⁽¹⁰⁾ and independently by Galasso *et al.*⁽²⁰⁾ Both workers originally could not account for the extra lines which most of the X-ray patterns of these compounds contained. Once Galasso *et al.*⁽²¹⁾ found that one of the compounds, Ba(Sr_{0.33}Ta_{0.67})O₃, had a new ordered perovskite structure; subsequent studies showed that many of the A²⁺(B_{0.33}D_{0.67})O₃ adopted this structure. The structure of Ba(Sr_{0.33}Ta_{0.67})O₃ is based on space group P3m1 with atoms in the unit cell at the following locations:

Ú

The structure is shown in Figs. 2.6 and 2.7. Note that if the perovskite structure is described as close-packed layers of A and oxygen ions perpendicular to the [111] direction with small B ions in the octahedral holes between the layers, then these B ions Sr and Ta each form planes of atoms. These planes are parallel to the close-packed layers and, since there are twice as many tantalum ions as strontium ions, the repeat scheme is two layers of tantalum ions and one of strontium ions. It is interesting that the ordered structure of A(B', sB'o, 5)O₃ type compounds when observed in the same way in the [111]

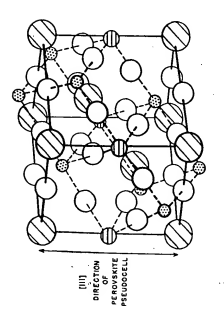




Fig. 2.6. The Ba(Sr_{0.33}Ta_{0.67})O₈ structure (after Galasso et al.⁽¹¹¹⁾).

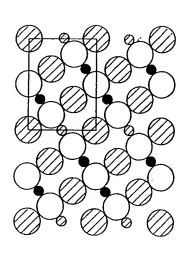




Fig. 2.7. Three-layer repeat sequence in Ba(SrossTao.67)Os.

The (110) plane (after Galasso et al.(11)

direction, consists of alternating layers, one of which contain

STRUCTURE OF PEROVSKITE-TYPE COMPOUNDS

B' ions and the other B" ions.

In some cases the ordering lines in the X-ray patterns were pounds which might be expected to be ordered because of A study of $Ba(B_{0.33}^2Ta_{0.67}^{5+})O_3$ compounds showed a decrease in the ordering as the difference in the size of the B2+ and Ta5+ ions became smaller. (22) The results were confirmed in a later investigation of $Ba(B_{0.33}^2Nb_{0.67}^{5+})O_3$ compounds.⁽²³⁾ very broad and became sharp when the samples were annealed at high temperatures. This observation was attributed to the existence of small ordering domains which grew at higher temperatures. Thus the X-ray patterns of some of the comthe large size difference of the B' and B' ions did not show the ordering lines if they could not be annealed at high temperatures because of low melting points.

Oxides of the $A^{2+}(B_{0.5}^{3+}B_{0.5}^{5+})O_3$, $A^{2+}(B_{0.5}^{2+}B_{0.5}^{6+})O_3$, $A^{2+}(B_{0.5}^{1+}B_{0.5}^{7+})O_3$ and $A^{3+}(B_{0.5}^{2+}B_{0.5}^{4+})O_3$ Types

The largest group of complex perovskite type compounds they adopt the structure shown in Fig. 2.5. It was postulated by Galasso et al. (20) that an ordered distribution of the B ions has the general formula A(BosBos)O3. When the strucis most probable when a large difference existed in either ture of these compounds is ordered, and most of them are, their charges or ionic radii. This hypothesis can be validated qualitatively by looking at the data below in Table 2.1.

Table 2.1. Structural Data for A(BxBy')O3 Type Compounds

Arrangement of B ions	Random ¹⁰ Ordered ¹⁴ Ordered ²⁴ Ordered ²¹ Random ¹⁰
Difference in ionic radii of B ions	0.04 0.05 0.46 0.44 0.05
Difference in charge of B ions	ପ୍ୟପ୍ତତ
Compound	Ba(Fe _{3.5} Ta ₆ [‡])O ₃ Ba(Mg _{6.5} ² W _{6.5})O ₃ Ba(La _{3.5} Ta _{6.5})O ₃ Ba(Sr _{6.5} ² 3 Ta _{6.5})O ₃ Ba(Zr _{6.5} ² 3 Ta _{6.5})O ₃ Ba(Zr _{6.5} ² 3 Nb _{6.5})O ₃

16

A study of $Ba(M_0^3 + Nb_0^5 + N)O_3^{(24)}$ type compounds indicated that the critical percentage difference in ionic radii between B ions which causes ordering lies between 7% and

ber of alkaline earth tungstates and molybdates of the A(A_{0.5}B_{0.5})O₃ type where A is an alkaline earth ion and B" is Mo or W have this structure. The structures of $\mathrm{Ba}(\mathrm{Ca}_{0.5}\mathrm{W}_{0.6})\mathrm{O}_3$ and became cubic only after heating to 500°C. Fresia et al. (15) found that other ions such as Zn²+, Fe²+, Co²+ and Ni²+ could be used as the B' ion in the ordered perovskite structure without distortion. However, when compounds were prepared with strontium in the A position the structures were distorted. It should be noted at this point that some authors index the tetragonal and orthorhombic distortion of the cubic ordered unit cell by using the ~ 8 Å edges, while others use an \sim 5.7 Å face diagonal for two axes and \sim 8 Å perature while the Ba(Sro, Wo.s)O3 structure was distorted and Ba(Ca_{0.5}Mo_{0.5})O₃ were reported to be cubic at room tem-The first compounds with this ordered structure were reported by Steward and Rooksby, (16) who found that a num edge for the third.

Sleight and Ward⁽¹⁷⁾ in a study of A(B²₆L⁰_{0,6})O₃ type perovskites also found that all of the compounds containing strontium in the A position had distorted structures. The unit cells were pseudomonoclinic, but the powder patterns were indexed on a smaller orthorhombic unit cell. Using a tolerance factor

$$E = \hat{K}_A + R_0/\sqrt{2} [(R_B^{2+} + R_B^{5+})/2 + R_0]$$

they calculated that a number of compounds containing barium in the A position should form the cubic ordered perovskite structure, and observed that they did form this structure.

Some of the most interesting compounds were those containing Mo⁵⁺, W⁵⁺ or Re⁵⁺ as one of the B ions and another paramagnetic ion as the other because of the ferromagnetic properties they exhibited. These compounds will be discussed in detail in another section.

By adjustment of the oxidation state by valence compensation the compounds $\text{Bu}(\text{Li}_{0.5}\text{Os}_{0.5})\text{O}_3$ and $\text{Ba}(\text{Na}_{0.5}\text{Os}_{0.5})\text{O}_3$

were prepared. (48) It appears that the osmium has a valence state of 7 in these compounds, a value not reported previously for this element. Similar compounds also were prepared with Dot the R. ion

Re'+ as the B" ion.

Lanthanum also has been introduced in the A position of the perovskite structure with two ions in the B position. The symmetry of the structure of these compounds has not been resolved, but it appears that they adopt some modification of the ordered perovskite structure. Preliminary single crystal studies indicate that they could have ordered cubic unit cells. (28) This would be interesting in light of the fact that there are no A³⁺B³⁺O₃ type ternary oxides with the simple cubic perovskite structure.

Oxides of the $A^{2+}(B_{0.25}^{1+}B_{0.75}^{5+})O_3$ Type

It was expected, because of the large charge difference in the B ions, that Ba(Na_{0.25}Ta_{0.75})O₃ and Sr(Na_{0.25}Ta_{0.75})O₃ would be ordered. However, no superstructure lines were observed in the X-ray patterns of these two compounds. (27) It should be noted that attempts to prepare compounds with smaller amounts of the B ion than 0.25 were not succeed.

Oxygen-deficient Oxides of the $A^{2+}(B_{0.5}^3B_{0.5}^4)O_{2.75}$ and $A^{2+}(B_{0.5}^2B_{0.5}^4)O_{2.75}$ Types

All compounds with the general formula A²⁺(B_{0,5}B_{0,5})O_{2,78} were found to have the ordered structure. Compounds containing tantalum as the B" ions were first reported by Brixner. (24) Later uranium and molybdenum were also found to form these oxygen-deficient compounds. (17, 28)

A summary of the oxides with the perovskite structure and structural data are presented in Table 2.2.

Compounds
Perovskite-type
for
Data ,
Cell
Unit
2.2.
TABLE

Compound	a (A)	A1+	$A^{1+B\delta+O_g}$	Remarks	Refer-	
	(1)		(1)		ences	
AgNbO3	7.888	15.660 7.888	7.888	$\beta = 90.57^{\circ}$	29	
AgTaO,	3.931	3.914	3.914 3.931	$\beta = 90.35^{\circ}$. 58	
CsIO3	9.324 or			cubio	30, 31	
KIO,	8.92			cubic	4, 31,	
	4.410 J OF			$\alpha = 80^{\circ}24'$ rhom-	32, 33,	
				bohedral	34	
KNb0,	3.9714	5.6946	5.6946 5.7203	orthorhombic	35	
KTaO3	3.9885			cubic	31, 36,	
					37	
NaNbo,	5.512	5.577	3.885	orthorhombic	35, 38	
NaTaO,	3.8851	5.4778	5.4778 6.6239	orthorhombic	39	
RbIO3	4.52 or			cubic	2, 31	
TIIO3	4.510			$\alpha = 89.34^{\circ} \text{ rhom}$	4, 40.	
•				bohedral	41	
		A3+]	A*+B4+0,			
		_				

		4	80 . TO . W		
BaCeO,	4.397				4, 5, 35,
BaFeO ₃	3.98		4.01	tetragonal	43, 44
BaMoO,	4.0404			•	45
BaPbO,	4.273				46
$BaPrO_3$	8.708 or				31, 47
BaPuO ₃	4.39				48
BaSnO ₃	4.117				1, 4, 5
BaThO,	4.480			monoelinic	4, 35,
	8.985				42, 47
BaTiO,	3.989	•	4.029	4.029 tetragonal	5
$BaUO_3$	4.387			pseudocubic	49
BaZrO ₃	4.192			cubic	ō
CaCeO ₃	7.70			cubic	31, 50
CaHfO				orthorhombic	51
CaMnO ₃	10.683	7.449	7.449 10.476	orthorhombic	13, 14,
				•	52
CaMoO,	7.80	7.77	7.80	$\beta = 91^{\circ}23'$ mono-	
				clinic	53
			-		

TABLE 2.2 (cont.)

Š	
_	
Ċ	١
.π÷	
α	
+	
◂	

Compound					Ļ
	a (A)	b (Å)	c (Å)	Remarks	Keter- ences
CaSnO	6.518	7.884	5.664	orthorhombic	4, 54
CaThO	8.74			monoclinic,	
	,			pseudocubic	31
CaTiO,	5.381	7.645		orthorhombic	54
CaUO,	6.78	8.29		orthorhombic	49
$CaVO_s$	5.326	7.547	5.352	orthorhombio	52
CaZrO	5.587	8.008		orthorhombic	20
CdCeO ₃	7.65			orthorhombic	
(1	1		or cubic	31, 60
CashO	0.047	0.677	7.867	orthorhombic	ء د
Cario	70. 1# 20.1	7 606	617	pseudocubic	, v
CdZrO,	1	20.		orthorhombic	٠ .
EuTio,	3.897				200
MgCeO,	8.54				31
PbCeO,	7.62			orthorhombic	31, 50
PbHf0,				pseudotetragonal	
PbSnO,	7.86		8.13	tetragonal	31
PbTiO,	3.896		4.136	tetragonal	89
PbZrO ₃	9.28			pseudocubic,	
				orthorhombic	
SrCeO ₃	5.986	8.531	6.125	orthorhombic	4, 5, 31
SrCoO.	7.725				<u> </u>
$SrFeO_3$	3.869				13
SrHfO ₃	4.069 or			orthorhombic	31
SrMoO.	3.9751				45
SrPbO.	5.864	5,949	8.336	orthorhombic	46
SrRuO,) 1)	cubic	69
SrSnO.	4.0334 or			cubic	1. 4. 60
° (_			:	•
SrThO,	48.8			pseudocubic	31
SrIIO3	3.904	6		ongno	۰,
aroo ₃	0.01	0°.0).T.0	orthornombic	A 4
SrZrO ₃	5.792 or	8.189	5.818	orthorhombic	4, 5, 31
				2000	_
		A ⁸ +	A*+B3+O3		
BiAlO,	7.61		7.94	tetragonal	31

PREPARATION OF PEROVSKITE-TYPE COMPOUNDS

(cont.)
2.5
TABLE

TABLE Z.Z (Cont.)		3+B3+C	A3+B3+O3 (cont.)	ut.)		
Compound	a (A)	b (Å)	c (Å)	Remarks	Refer- ences	
BiCrO3	3.90	3.87	3.90	$\alpha = \gamma = 90^{\circ}35'$ triolinic $\beta = 80^{\circ}10'$	61	
BiMnO ₃	3.93	3.98	3.93	$\alpha = \gamma = 91^{\circ}25'$ triclinic	61	
CeAlO, CeCrO,	3.767 3.866		3.794	tetragonal	62, 63 63, 64, 65	
CeFeO	3.900			pseudocubic,	64. 63	
CeGaO3	3.879			oubio,		
CeScO	(orthorhombic	64	
CeVO.	3.90		8.08	tetragonal	90, 00 31	
DyAlo,	6.21	5.31	7.40	orthorhombic	67	-
DykeO.	3.70	9.0	70.1	cubic	89	
EuAlO,	6.271	6.292	7.458		63, 69	
EuCrO, EuFeO,	3.803 6.371	5.611	7.686	Ğ	63 6, 63,	
${ m FeBiO_3}$	7.64 3.966} or			$\alpha = 89^{\circ}28'$ rhombohedral	69 70, 71,	
GdAlO3	5.247	5.304	7.447	GdFeO ₃ structure	63, 67,	
GdCoO3 GdCrO3	3.732 5.312	3.807	3.676	9 9	73	
GdFeO ₃ GdMnO ₃	5.346 3.82	5.616			6, 75 14	
GdScO	5.487	5.756	7.925		74	
GdVO3	5.345	5.623	7.638	Ğ	74	
LaAlO ₃	3.788			$\alpha = 90^{\circ}4' \text{ rhom-}$ bohedral	ıo.	

TABLE 2.2 (cont.)

TABLE 2.2 (cont.)	t.)	A8+	A8+B3+O3	(cont.)	
Compound	a (Å)	b (Å)	c (Å)	Remarks	Refer- ences
LaCoO3	3.824 7.661}or			$\alpha = 90^{\circ}42'$ rhombohedral	13, 63, 65, 76,
LaCrO	5.477	5.514	7.755	GdFeO ₃	! • i
LaFeO ₃	5.556	5.565	7.862	structure GdFeO ₃	74
LaGaO,	5.496	5.524	7.787	structure GdFeO,	6, 6
LaInO,	5.723	8.207	5.914	structure orthorhombic	74 5,64
LaNio	7.676			$\alpha = 90^{\circ}41'$	
LaRhO ₃ LaScO ₃	3.94	5.787	8:098	pseudocubic GdFeO ₃	76, 80
i i	(structure	
LaTiO, LaVO,	$\frac{3.92}{3.99}$			cubic	81, 82 65, 66, 83
LaYOs	3.752			orthorhombic rhombohedral	84 63. 67.
, 1)				
NdCoO.	3.777	707	7 605	CAR _e O	65, 73
INGCE C3	714.0	¥.0	060.	structure	74
NdFeO3	5.441	6.573	7.753	GdFeO	6 64
NdGaO3	5.426	5.502	7.706	GdFeO	5
				structure	74
NdInO ₃ NdMnO ₃	5.627 3.80	8.121	5.891	orthorhombic	68 68
NdScO	5.574	5.771	7.998	GdFeO	
OAPN	5.440	5.589	7.733	structure GdFeO.	64, 74
, , , , , , , , , , , , , , , , , , ,)		structure	
PrA10,	$\frac{3.757}{5.31}$ or			$\alpha = 60^{\circ}20'$	48, 53, 69
PrCoO3	3.787			$\alpha = 90^{\circ}13'$	
0,0	777	707	7 710	rhombohedral	65, 73
, TOTO 3	# # * * * * * * * * * * * * * * * * * *				74

TABLE 2.2 (cont.)

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TABLE 2.2 (cont.)

A3+B3+O3 (cont.)

Compound	a (Å)	b (Å)	c (Å)	Remarks	Refer- ences
PrFeO.	5.496	5.578	7.810	GdFeO,	
79 1				structure	74
PrGaO3	5.465	5.495	7.729	GareOs	74
PrMnO.	3.82				89
PrScO,	5.615	5.776	8.027	GdFeO,	
•					74
PrVO ₃	5.477	5.545	7.759	Ğ	
' '				structure	74
PuAlO ₃	6.33			$\alpha = 50 4$	8
DuCrO.	5.46	5.51	7.76	GdFeO.	
, , , , , , , , , , , , , , , , , , ,				structure	48
PuMnO.	3.86			pseudocubic	48
PuVO,	6.48	5.61	7.78	GdFeO,	
	_			structure	48
SmA10,	5.285	5.290	7.473	GdFeO,	
,				structure	
-					
SmCoO,	3.747	3.803			66, 73
SmCrO,	5.372	5.502	7.650	GdFeO,	
•				structure	7.4
SmFeO,	5.394	5.692	7.711	ŏ	,
•				structure	6, 64
SmInO	5.589	8.082	5.886	orthorhombic	δ, Σ
SmVO3	3.89	1		Caro	3
YAIO3	6.179	6.329	0,5.7		9
,	t .		2 2 2	Č	1
x CrO3	0.24 (0.00			6, 7,
$ m YFeO_{s}$	5.302	6.589	7.622	GdFeO,	
					ĝ
YScO,	5.431	5.712	7.894	GdFeO ₃	74

•	orthorhombic orthorhombic
ArBO, and ABO,-	7.86
30, an	3.91
A.F	3.89 3.90
	Ce.33NbO2 Ce.33TaO3

22

Perferance ence ence ence ence ence ence ence	TABLE 2.2 (cont.)	A.BO, and ABO,	1 ABO	-= (cont.)	, (: ₁	
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	Compound	a (Å)	b (Å)	c (Å)	Remarks	Refer- ences
a. 3.87 3.89 7.73 orthorhombic 10 3.91 7.88 tetragonal 10 10 10 3.91 7.76 orthorhombic 10 10 3.91 7.77 tetragonal 10 10 3.91 7.77 tetragonal 10 3.92 7.77 orthorhombic 10 3.89 7.77 orthorhombic 10 3.89 7.77 orthorhombic 10 3.89 7.75 tetragonal 10 3.82 7.78 orthorhombic 10 11.068 7.505 5.378 orthorhombic 85 ($x=1$) $x=0.2$ orthorhombic $x=0.2$ $x=0.3$ orthorhombic $x=0.3$	Dyo.33TaOs	3.83	3.83	7.75		9
$\begin{array}{cccccccccccccccccccccccccccccccccccc$					monoclinic	2 :
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	Gd, "TaO,	3.87	3.89	7.73	orthorhombic	01
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	La, 1,NbO	3.91		7.90	tetragonal	01
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	Lac 21TaO	3.92		7.88	tetragonal	2
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	Nd. "Nbo.	3.90	3.91	7.76	orthorhombic	10
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	Nd Tab.	3.91		7.77	tetragonal	10
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	Dr. NBO	3.91	3.92	7.77	orthorhombic	10
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	D. T.O.	3.91	3.92	7.78	orthorhombic	10
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	CT To To C	3,83		7.75	tetragonal	10
monoclinic 10 $(x=1)$ 3.79 3.80 7.70 $(x=91.6^{\circ})$ monoclinic 10 $(x=1)$ 3.72 $(x=1)$ cubic $(x=1)$ 3.8622 $(x=0.2)$ cubic $(x=0.45)$ cubic $(x=0.45)$ cubic $(x=0.7-0.9)$ $(x=0.45)$ $(x=0.45)$ $(x=0.46)$ $(x$	V. TaO	3.82	3.83	7.74	ر 190.9° م	
$Nb_{1-2x}^{4-2x}O_{s} \begin{pmatrix} x=0.2 \\ x=1 \end{pmatrix}$ $Nb_{1-2x}^{4-2x}O_{s} \begin{pmatrix} x=0.2 \\ x=0.35 \end{pmatrix}$ $Nb_{1-2x}^{4-2x}O_{s} \begin{pmatrix} x=0.2 \\ x=0.2 \end{pmatrix}$ $A(B_{0,67}B_{0,33}^{\alpha})O_{s}$ $A(B_{0,67}B_{0,33}^{\alpha})O_{s}$ $NH_{0,83}^{4-3}O_{s} \begin{pmatrix} x=0.2 \\ x=0.7-0.9 \\ x=0.7-0.9 \end{pmatrix}$ $A(B_{0,67}B_{0,33}^{\alpha})O_{s}$ $A(B_{0,67}B_{0,33}^{\alpha})O_{s}$ $A(B_{0,67}B_{0,33}^{\alpha})O_{s}$ $A(B_{0,67}B_{0,33}^{\alpha})O_{s}$ $A(B_{0,67}B_{0,33}^{\alpha})O_{s}$ $A(B_{0,67}B_{0,33}^{\alpha})O_{s}$	50.83 + C. 8.0 +					10
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	Yb,,33TaO,	3.79	3.80	7.70		
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$		•	1		monoclinic	01
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	Cao.sTaOs	11.068	7.505		orthornombic	Q X
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	Li, WO,	(x=1)			:	
$\begin{array}{cccccccccccccccccccccccccccccccccccc$		3.72				
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$		(1 -1)			l	
$\begin{array}{c} x = 0.7 - 1.0 & 8, \\ (x = 0.2) & 0.00 & 0.00 \\ 3.981 & 0.00 & 0.00 \\ 4.016 & 0.00 & 0.00 & 0.00 \\ 4.016 & 0.00 & 0.00 & 0.00 \\ 4.016 & 0.00 & 0.00 & 0.00 \\ A & B & B & B & B & B \\ & & & & & & & & &$	Na. WCs	(X = 1)	_		cubic	
						8, 50,
						87, 88,
		-				89,80
$ \begin{array}{c} (x=0.2) \\ 3.981 \\ (x=0.45) \\ 4.016 \end{array} $ $ \begin{array}{c} (x=0.7-0.9 \\ (x=0.7-0.9) \\ (x=0.$						91, 92
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	Sro.5+2Nb22 Nb1+22O8				, it	σ
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$			_		capic	·
A(B' _{0.93})O ₈ A(B' _{0.97} B' _{0.33})O ₈ A(B' _{0.97} B' _{0.33})O ₈ A(B' _{0.97} B' _{0.33})O ₈ B(NH ₄) ₈ FeF ₆ O(NH ₄) ₈ FeF ₆ O(NH ₄) ₈ FeF ₆		4.016				
A(B _{0.67} B _{0.33})O ₃ B(NH ₄) ₈ FeF ₆ O(NH ₄) ₈ FeF ₆ O(NH ₄) ₈ FeF ₆	CaMnO _{1.}					13, 14
A(B' _{0.97} B' _{0.33})O ₃ A(B' _{0.97} B' _{0.33})O ₃ A(B' _{0.938})O ₃ A(B' _{0.938})O ₃ B.386 A(B' _{0.97} B' _{0.33})O ₃ B.386	STOOL ST					13
A(B _{0.67} B _{0.33})O ₂ A(B _{0.67} B _{0.33})O ₂ (NH ₄) ₂ FeF ₆ (NH ₄) ₂ FeF ₆	SrFeO					11, 12,
A(B' _{0.33})O ₃ A(B' _{0.33})O ₃ A(B _{0.33})O ₃ A(B _{0.33})O ₃ A(B _{0.33})O ₃	1 000					13
A(B _{0,67} B _{0,33})O ₃ W _{0,33})O ₃ W _{0,33})O ₃ 8.386 (NH ₄) ₈ FeF ₆	SrTiO3_2					83
A(B _{0.67} B _{0.33})O ₃ 8.386 (NH ₄) ₂ FeF ₆	SrVO3-2		_			73 XO
A(B _{0.67} B _{0.33})O ₃ 8.386 (NH ₄) ₂ FeF ₆			_	_		_
8.386 (NH ₄) ₈ FeF ₆		A((Bí, 1, B″	.33)08		
8.386 (NH _{4,9} FeF ₆						8
8.386 (IN IL) SEE OF SEE	Ba(Alo. 67 Wo. 33) Os				T. T. T. T.	3
	Ba(Dyo.e7 Wo.33)U3	8.380			B TH TR (PT NI)	69

PREPARATION OF PEROVSKITE-TYPE COMPOUNDS

TABLE 2.2 (cont.)

A(B'0.87B'0.83)O₃ (cont.)

Ba(Er _{0.67} W _{0.33})O ₃ . 8 Ba(Eu _{0.67} W _{0.33})O ₂ 8 Ba(Fe _{0.67} U _{0.33})O ₃ 8 Ba(Gd _{0.67} W _{0.33})O ₃ 8	8.386	-			
	-			(NH,),FeF,	3
	8.605			structure (NH ₄) ₃ FeF ₆	
	8.232			structure (NH,),FeF,	83
	-			structure	17
-	8.411		•	(NH4)3FeF6 structure	93 94
Ba(In _{0.67} U _{0.33})O ₃ 8	8.512			(NH,)3FeF,	
Ba(In _{0.67} , W _{0.33})O ₃ 8	8.321			structure ordered	17
Ba(La _{0.67} W _{0.33})O ₃ 8	8.58			structure (NH ₄) ₃ FeF ₆	63
!				structure	94
Ba(Lu _{0.67} W _{0.33})O ₃ Ba(Nd _{0.67} W _{0.33})O ₃ 8	8.513	-		(NH,)3FeF	93
Ba(Sc _{0.67} U _{0.33})O ₃ 8	8.49			structure (NH ₄) ₃ FeF ₆	£6 5
Ba(Sc. 67 Wo. 33) O3 8	8.24	•		structure (NH ₄) ₃ FeF ₆	17
	8.70		-	structure (NH1),FeF,	15, 93
	i c			structure	17
Ba(Y _{0.67} W _{0.33})O ₃ 8	8.374			$(NH_4)_3$ FeF $_6$	63
Ba(Yb _{0.6} , W _{0.33})O ₃ Pb(Fe _{0.52} W _{0.33})O ₂		•			93
	8.01		-	(NH4)3FeF	6
$Sr(Cr_{0.87}U_{0.33})O_3$ 8	8.00			structure . (NH4)3FeFs	 8
	68			structure (NH.), FeF.	17
) }		<u> </u>	structure	18
Sr(Fe _{0.67} W _{0.33})O ₃ 3 Sr(In, c,Re _{0.33})O ₃ 8	3.945		3.951	tetragonal (NH ₄) ₂ FeF	47, 94
	. 0	u u	9	structure	18
La(Co. 67 Sbo. 33) O3 5	5.57	5.57	7.89	orthorhombic	94
•	····				

TABLE 2.2 (cont.)

A2+(B2+3B5+,)O3 (cont.)

Compound	a (Å)	b (Å)	c (Å)	Remarks	Refer- ences
Ba(Ca _{0.33} Nb _{0.67})O ₃	6.92		7.25	hex. ordered Ba(Sr _{0.83} Ta _{0.67})O _{:1}	
Ba(Ca _{0.33} Ta _{0.67})O ₃	5.895		7.284	structure 7.284 hex. ordered Ba(Sr _{0.83} Ta _{0.67})O ₃	23
Ba(Cd _{0.33} Nb _{0.67})O ₃ Ba(Cd _{0.33} Ta _{0.67})O ₅	4.168			structure	21, 97, 98. 23 22
Ba(Co.33Taois7)Os	5.776		7.082	7.082 hex. ordered Ba(Sr _{0.33} Ta _{0.67})O ₃	
Ba(Cu.33Nb.67)O3 Ba(Fe.33Nb.67)O3 Ba(Fe.77	8.04		8.40	structure tetragonal	19, 22 94 23 20
Ba(Mg _{0.33} Nb _{0.67})O ₃	6.77	-	7.08	hex. ordered Ba(Sr _{0.88} Ta _{0.67})O ₃	3
Ba(Mg _{0.33} Ta _{0.67})O ₃	5.782		7.067	structure 7.067 hex. ordered Ba(Sr _{0.89} Ta _{0.67})O ₈	zu, zs, 94-
Ba(Mn _{0.33} Nb _{0.67})O ₃ Ba(Mn _{0.33} Ta _{0.67})O ₃	5.819		7.127	atructure 7.127 hex. ordered Ba(SrTa)O.	22, 98 93
Ba(Ni _{0.33} Nb _{0.67})O ₃ Ba(Ni _{0.33} Ta _{0.67})O ₃	4.074		7.052	structure 87.052 hex. ordered	22 19, 23, 96
Ba(Pb _{0.33} Nb _{0.67})O ₃ Ba(Pb _{0.33} Ta _{0.67})O ₃	4.26			Ba(Sr _{0.33} Ta _{0.67})O ₈ structure	19, 22, 98 23 22
Ba(Sr _{0.33} Ta _{0.67})O ₃	5.95		7.47	hex. ordered Ba(Sr _{0.33} Ta _{0.67})O ₈ structure	20, 21
Ba(Zn _{0.33} N b _{0.67})O ₈ Ba(Zn _{0.33} Ta _{0.67})O ₃	4.094 5.782		7.097	7.097 hex. ordered Ba(Sr _{0.83} Ta _{0.67})O ₈ structure	20, 23, 90
Ca(Ni _{0.33} Nb _{0.67})O ₃ Ca(Ni _{0.33} Ta _{0.67})O ₃ Pb(Co _{0.33} Nb _{0.67})O ₃ Pb(Co _{0.33} Ta _{0.67})O ₃	3.88 3.93 4.04 4.01				96 19 96, 99 99

A2+(B2+85+7)O3 (cont.) TABLE 2.2 (cont.)

Compound	a (A)	b (A)	c (Å)	Remarks	Refer- ences
Pb(Mg _{0.33} Nb _{0.67})O ₃ Pb(Mg _{0.33} Ta _{0.67})O ₃	4.041				96, 100 96, 99
Pb(Mi _{0.33} Nb _{0.67})O ₃ Pb(Ni _{0.33} Nb _{0.67})O ₃					
Pb(N10.33 Ta0.67) U3 Pb(Zn, 33 Nb, 27) O3	4.01				96, 99 90
Sr(Ca _{0.33} Nb _{0.67})O ₃			7.16	hex. ordered))
Sr(Ca _{0.33} Sb _{0.87})O ₃	8.17			Da(Sr. 83 L B _{0.67}) U ₃ structure (NH ₄),FeF,	23
Sr(Ca _{0.13} Ta _{0.67})O ₃	5.764		7.096	structure 7.096 hex. ordered	94
				Ba(Sro.83 Tao.67)Og	66
Sr(Cd _{0.83} Nb _{0.67})O ₈ Sr(Co. 22Nb _{2.23})O ₃	4.089			NH) Ref	53
Sr(Co Sb V)	1 00			Structure	94
Ot (CO ₀ .835 C ₀ .67) C ₃				Structure	94
Sr(Co _{0.33} Ta _{0.67})O ₃	5.630		6.937	6.937 hex. ordered	!
				Ba(Sr _{0.88} Ta _{0.67})O ₈ structure	20. 22
Sr(Cu _{0.33} Sb _{0.87})O ₃	7.84		8.19	tetragonal	
Sr(Mg, 83N Do, 87) O3	5.88		6.98	4.013 tetragonal 6.98 hex. ordered	27
		-		Ba(Sro.83 Teo.67)Os	
Sr(Mg _{0.83} Sb _{0.67})O ₃	7.96			structure (NH ₄) ₈ FeF ₆	23
Sr(Mg, ",Ta,",)O,	5.652		6.951	structure	94
				Ba(Sr _{0.83} Ta _{0.67})O ₃	
Sr(Mn _{0.33} Nb _{0.67})O ₃				structure	19, 22 93 03
Sr(Ni _{0.83} Nb _{0.67})O ₃	5.64		6.90	hex. ordered	9
				Ba(Sro.33Tao.67)O3	93 06
Sr(Ni _{0.33} Ta _{0.67})O ₃	5.607		6.923	6.923 hex. ordered	
	-			Ba(Sr _{0.88} Ta _{0.67})U ₈ structure	20, 22
Sr(Pb _{0.33} Nb _{0.67})O ₃ Sr(Pb _{0.33} Ta _{0.67})O ₃					93
			-		-

TAB

A2+(B2+3B6+7)O3 (cont.)

STRUCTURE OF PEROVSKITE-TYPE COMPOUNDS

cont.)	
2.2	•
BLE	

Compound	a (Å)	b (Å)	c (Å)	Remarks	Refer- ences
Sr(Zno. 33Nbo. 67)O3	5.66		6.95	hez. ordered Ba(Sr, s.Ta, g.)O.	
$\mathrm{Sr}(\mathrm{Zn_{0.33}Ta_{0.67}})\mathrm{O_3}$	5.664		6.951	6.951 hex. ordered	20, 23
				Ba(Sr _{0.83} Ta _{0.67})O ₃ structure	20, 22
		A*+(B3	A*+(B3;B3;5)O3)3	
Ba(Bio.sNbo.s)Os	8.630			(NH,)3FeF	
Ba(Bio., Tao., 5)Os	8.568			structure (NH4) ₃ FeF ₀	101
Ba(Ce, Nb, 10,	4.293	_	•	structure	101
$Ba(Ce_{0.\delta}Pa_{0.\delta})O_{\delta}$	8.800			(NH,)3FeFe	
Ba(Coo.gNbo.s)Os	4.06			structure (NH ₄),FeF,	103
Ba(Co, Re, 1)O,	8.086			structure (NH.),FeF.	9 4
		•		structure	18
Ba(Cro., Wo., s)O, The Cro., W. VO	900		19 0	1000000+0+	104
Ba(Dy,,,Nb,,,)O,	8.437		70.0	(NH,),FeF,	# 0
, d	i c			structure	25, 102
Ba(Dy _{0.5} Fa _{0.5})O ₃	8.740		•	(NH4)3KeFg	103
Ba(Dyo., Tao., 8)O,	8.545			(NH4)3FeF	2
Ba(Er. Nb)O.	8.427			structure (NH.), FeF.	106
8) /9:00:0				structure	25, 102
Ba(Er _{0.5} Pa _{0.5})O ₃	8.716			(NH4) ₃ FeF ₆	103
Ba(Er _{0.8} Re _{0.8})O ₈	8.354			(NH,)3FeF	
Ba(Ero, Tao, 1)O3	8.423			etructure (NH,),FeF,	8
O, 11 -0/-0	0			structure	105
D&(Ed. 0.8 U.8 J U.8	0.0		-	Structure	104
Ba(Eu., Nb,, 6)03	8.507			(NH,)3FeF	
Ba(Eu., Pa,,)O,	8.783			Structure (NH4) ₃ FeF	20, 102
				structure	103

A2+(B3+B5+)O3 (cont.) TABLE 2.2 (cont.)

									•								•						••			
Refer- ences	105	28 20, 25, 106	0 -	20, 103	25, 102	103	č	}	94	105	26, 102	103		901	25, 107	18		103	18	94, 108	. ;	e 6	104	25, 102, 106	301	ent
Remarks	(NH,)3FeFe	structure	(NH,) FeF.	Burucume (NH) FeF	structure	structure	(NH4)3FeFg	(NH4),FeF	structure	tetragonal (NH ₆) ₃ FeF ₆	structure	structure	(NH4)3FeF	structure (NH ₄) ₃ FeF,	structure (NH.), FaF	structure	(NH4)3FeF	structure (NH4),FeF,	structure	structure	(NH4) ₈ FeF _{6,}	structure (NH,),FeF,	structure	tetragonal	(NH,)3FeF	structure
c (Å)							<u></u> .			8.513				<u> </u>						-				8.690		
b (Å)																				•						
a (Å)	8.506 8.08	4.06	8.05	4.056 8.496	, t	9.1.14	8.431	8.44		8.487	0 2 7 3 0	0.5	8.442	8.279	8 224		8.596	8.258	9 960	604.0	8.280	8.52	1	8.607	8.885	
Compound	Ba(Eu _{0.8} Ta _{0.8})O ₈ Ba(Fe _{0.8} Mo _{0.6})O ₈	Ba(Fe, Nb, s)O3	Ba(Fe _{0.8} Re _{0.8})O ₃	Ba(Fe _{0.8} Ta _{0.8})O ₃	B.(Gd B.)	Da(Juo. 8 F 20. 8) U3	Ba(Gd _{0.5} Re _{0.5})O ₂	Ba(Gdo, Sbo, 6)O3	Ç	Ba(Hd _{0.8} Tia _{0.8})O ₃ Ba(Ho _{0.8} Nb _{0.8})O ₃	Re/Ho De 10	La(1100.81 a _{0.8} /O ₃	Ba(Ho _{0.8} Ta _{0.8})O ₃	Ba(Ino.gNbo.g)O3	Ba(In, Os. 1)O.	0.6 (0.6) (3	Ba(In _{0.8} Pa _{0.5})O ₃	Ba(In, Re, s)O,	Ba(In Sh 10	La(110.8000.5)O3	Ba(In _{0.8} Ta _{0.8})O ₃	Ba(In, U, s)O,		La(La _{0.8} Nb _{0.8})O ₃	Ba(La _{0.8} Pa _{0.5})O ₃	

STRUCTURE OF PEROVSKITE-TYPE COMPOUNDS TABLE 2.2 (cont.)

 $A^{2+}(B_{0.5}^{3+}B_{0.5}^{5+})O_{3}$ (cont.)

Compound	a (Å)	b (Å)	c (A)	Remarks	Refer- ences
Ba(La _{0.5} Re _{0.5})O ₃	8.58			(NH4)3FeF	
$\mathrm{Ba}(\mathrm{La_{0.5}Ta_{0.5}})\mathrm{O_{3}}$	8.611	8.639	8.764	structure (NH ₄) ₃ FeF ₆	18
Ç,	6			en nonne	106
Ba(Lu _{0.5} N b _{0.5})O ₃	8.364			(NH4) ₃ FeF ₆ structure	25, 102
$\mathrm{Ba}(\mathrm{Lu}_{0.8}\mathrm{Pa}_{0.8})\mathrm{O}_3$	8.666			(NH,)3FeF	201
$Ba(Lu_{0.5}Ta_{0.5})O_3$	8.372			Structure (NH4)3FeF,	601
Ba(Mno.sNbo.s)O3	4.083		,	structure	106
Ba(Mno.sReo.s)O3	8.18			(NH,) ₃ FeF,	ø
$\mathrm{Ba}(\mathrm{Mn}_{\mathrm{0.6}}\mathrm{Ta}_{\mathrm{0.6}})\mathrm{O}_{\mathrm{s}}$	4.076				106
$Ba(Nd_{0.5}Nb_{0.5})O_3$	8.540			(NH ₄) ₃ FeF ₆	25, 102.
			-		106
$Ba(Nd_{0.8}Pa_{0.8})O_3$	8.840			(NH4),FeF	601
Ba(Nd, Re, 1)0,	8.51			Structure (NH4),FeF,	703
9 (8:0 8:0				structure	18
Ba(Nd _{0.8} Ta _{0.8})O ₃	8.556				105, 106
Ba(Pr., Nb.,)0,	4.27				102, 106
Ba(Pro.,Pao.,s)O3	8.862		-	$(NH_4)_3FeF_6$	103
$\mathrm{Ba}(\mathrm{Pr}_{a,s}\mathrm{Ta}_{a,s})\mathrm{O}_{s}$	4.27			structure (NH4)3FeF	108
Ba(Rho., Nbo.,)O3	8.17			structure	94
$\mathrm{Ba}(\mathrm{Kh}_{\mathfrak{d},\mathfrak{s}}\mathrm{U}_{\mathfrak{d},\mathfrak{s}})\mathrm{U}_{\mathfrak{s}}$			•	nexagonal BaTiO,	104
Ba(Sco., NDo., 5)O3	4.121			0.00 (DIV)	96, 102
58(5c _{0.8} 08 _{0.8})03	8.152			(IV FL 4) 3F SF 6 structure	18
$\mathrm{Ba}(\mathrm{Sc}_{\mathfrak{d}.\mathfrak{s}}\mathrm{Pa}_{\mathfrak{d}.\mathfrak{s}})\mathrm{O}_{\mathfrak{s}}$	8.549			$(NH_4)_3FeF_6$	103
$\mathrm{Ba}(\mathrm{Sc_{0.8}Re_{0.8}})\mathrm{O_{3}}$	8.163			(NH4)3FoF	3
Or 40 '0/'B	0 107			Structure	8
Da(DC ₀ , gO D ₀ , g) O ₃				structure	108
$\mathrm{Ba}(\mathrm{Sc}_{0.5}\mathrm{Ta}_{0.5})\mathrm{O}_{3}$	8.222			(NH4)3FeF6	95 66
				901 00000	20, 00

30

A*+(B3+B5+)O3 (cont.)

TABLE 2.2 (cont.)

A2+(B3+B	
₹	
cont.)	
2.2	
TABLE	
_	

Refer-

Remarks

c (Å)

b (Å)

a (Å)

Compound

ences

TABLE 2.2 (cont.)		A ²⁺ (B _{0.5} ³⁺ B _{0.5} ⁵)O ₃ (cont.)	;) o° (t	ont.)	
Compound	a(A)	b(A)	c(A)	Remarks	Refer- ences
Ca(Cr., Nb,, s)O3	3.85	3.85	3.85	$\beta = 90^{\circ}47'$	
				monoclinic	109
$Ca(Cr_0, Os_0, s)O_8$	5.38	7.66	5.47	orthorhombic	18
$Ca(Cr_0, Re_0, s)O_3$	5.38	7.67	5.47		18
$Ca(Cr_0, Ta_0, s)O_3$	3.85	3.85	3.85	$\beta = 90^{\circ}45'$	
				monoclinic	109
$C_{a}(C_{r_0,s}W_{0,s})O_{s}$	5.47	7.70	5.35	orthorhombic	28
Ca(L)yo.sN Do.s)U3	4.03	4.03	4.0	$\rho = 32.20$ monoclinic	109
$Ca(Dy_{0.8}Ta_{0.8})O_8$	4.03	4.03	4.03	$\beta = 92^{\circ}24'$	
· ;	9	3	90	monoclinic	103
Ca(Ero, giv Do. 8) C3	4.02	4.01	4.02	$p = .92 \cdot 11$	109
Ca(Er, Ta, 1)O,	4.02	4.01	4.02	$\beta = 92^{\circ}10'$) •
				monoclinic	109
$Ca(Fe_05Mo_05)O_3$	5.53	7.73	5.42		28
$Ca(Fe_0.sNb_0.s)O_3$	3.89	3.88	3.89	$\beta = 91^{\circ}2'$	
:			1	monoclinic	109
Ca(Fee. Sbo. s)O3	5.54	5.47	7.74		84
$\mathrm{Ca}(\mathrm{Fe}_{\mathfrak{d}.\mathfrak{s}}\mathrm{Ta}_{\mathfrak{d}.\mathfrak{s}})\mathrm{O}_{\mathfrak{s}}$	3.89	3.88	3.89	$\beta = 91^{\circ}7'$	
OX 118 FD/-0	50	3	60 1	monoclinic ,	601
CE (GG0.81N Do.8) C3	4.0o	4.0.4	30.4	p = 32 #2	100
Ca(Gd, Ta, 10,	4.03	4.04	4.05	$\beta = 92^{\circ}41'$	2
201000000000000000000000000000000000000					∶109
Ca(Ho,,Nb,,,)O3	4.02	4.02	4.02	$\beta = 92^{\circ}19'$	
!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!	:			Ĕ	109
Ca(Ho, , Ta, ,)O3	4.03	4.02	4.03	$\beta = 92^{\circ}16^{\circ}$	001
Ca(InNb)O.	3.97	3.95	3.97	$\beta = 91^{\circ}53'$	2
8 1000000					109
$Ca(In_{0.8}Ta_{0.8})O_{8}$	3.97	3.96	3.97	$\beta = 91^{\circ}51'$	
		į			108
$Ca(La_{o.s}Nb_{o.s})O_3$	4.07	4.07	4.07	$\beta = 92^{\circ}8'$	100
Ca(La, Ta,)O,	4.07	4.07	4.07	$\beta = 92^{\circ}9'$	9
					109
$\mathrm{Ca}(\mathrm{Mn}_{\mathfrak{o}.\mathfrak{s}}\mathrm{Ta}_{\mathfrak{o}.\mathfrak{s}})\mathrm{O}_{\mathfrak{s}}$	3.90	3.87	3.90	$\beta = 91^{\circ}9'$	109
Ca(Nd, Nb, 1)0,	4.05	4.05	4.05	$\beta = 92^{\circ}28'$	
2) (9.0))	}	,		109

105 102, 106

(NH₄)₃FeF₈ structure

103

(NH,)3FeF (NH4)3FeF (NH,)3FeF, (NH4)3FeFe (NH4)3FeF

8.718

 $\mathrm{Ba}(\mathrm{Y}_{0.6}\mathrm{Nb}_{0.6})\mathrm{O_3}$ $\mathrm{Ba}(\mathrm{Y}_{0.6}\mathrm{Pa}_{0.6})\mathrm{O_3}$

8.372 8.433

Ba(Y0.8Req.8)O3

Ba(Yo.8Tao.5)O3 Ba(Yo.5U0.8)O3

structure

18

structure structure

25, 102

structure (NH4)₃FeF

 $(NH_4)_3FeF_6$

8.408 8.692 8.406 4.200

Ba(Tmo.sNbo.s)Os

Ba(Tm, Pa, 103 Ba(Tm_{0.8}Ta_{0.8})O₃

structure

103

structure

105, 106

104

structure structure

8.374

 $Ba(Yb_0._8Nb_0._8)O_3$

8.69

25, 96,

96, 105

structure $\beta = 90^{\circ}15'$

3.80 3.80

3.81 3.81

Ca(Al_{0.8}Nb_{0.8})O₃ $Ca(Al_{0.8}Ta_{0.8})O_3$

109

monoclinic $\beta = 90^{\circ}17'$ monoclinic

3.81 3.81

109 94 28

orthorhombic orthorhombic

7.73

5.43 7.70

6.60 5.49

Ca(Co, W, s)O3 Ca(Cr, Mo, s)O3

103 102

> NH,)3FeF. (NH,),FeF,

8.678 8.390

Ba(Ybo.,8Pao.,5)O3 Ba(Ybo. 5Tao. 5)03

structure

103 105, 106

structure

8.519 4.229

Ba(Sm_{0.8}Ta_{0.8})O₃ Ba(Tb_{0.8}Nb_{0.8})O₃ Ba(Tb_{0.8}Pa_{0.8})O₃

8.753

8.42

 $Ba(Tl_{0.5}Ta_{0.5})O_3$

(NH,)3FeF

8.792

Ba(Smo. 8Pao. 8)O3

103 108

(NH4)₃FeF₆ structure

(NH,)3FeF

25, 102, 106

104

(NH₄)₃FeF₆ structure

NH4)3FeF structure

8.518

Ba(Sm_{0.8}Nb_{0.8})O₃

Ba(Sco. & Uo. 5)O3

32

(cont.)
$A^{2+}(B_{0.5}^{3+}B_{0.5}^{6+})O_3$
(cont.)
TABLE 2.2 <i>(</i>

Ca(Ndo., Tao., s)O3 Ca(Nio., Wo., s)O3 Co(Nio., Wr., s)O3					ences	
0.8We.s)O3	4.05	4.05	4.05	$\beta = 92^{\circ}25'$		
2	5.65	5.40	7.70	monoclinic orthorhombic	109	
0.6-1-0.6/08	4.06	4.05	4.08	$\beta = 92^{\circ}25'$;	7
Ca(Pr _{0.8} Ta _{0.8})O ₃	4.06	4.05	4.06	monoclinic $\beta = 92^{\circ}22'$	109	
Carles 10	70	7 00	69	monoclinic	109	
Ca(Smo.,8Nbo.,8)O3	4.04	4.04	4.04	$\beta = 92^{\circ}42'$	07	
Ca(Sm _{0.8} Ta _{0.8})O ₃	4.05	4.04	4.05	monoclinic $\beta = 92^{\circ}28'$	109	
Ca(Tb, Nb, A)O,	4.03	4.03	4.03		109	•
Ca(Tb, Ta, 1)O.	4.03	4.03	4.03		109	•
י לו	ć	3			109	
Ca(± 0.514 D _{0.8}) C ₃	#.03	4.02	4.03	p = 92-23 monoclinie	109	
Ca(Y _{0.8} Ta _{0.8})O ₃	4.03	4.05	4.03	$\beta = 92^{\circ}23'$		
Ca(Yba, Nba, k)O,	4.01	4.00	4.01	monoclinic $\beta = 92^{\circ}0'$	109	
					109	
Ca(Y b _{0.8} Ta _{0.8})O ₃	4.01	4.00	4.01	$\beta = 92^{\circ}3'$	001	
Pb(Fe, Nb, 1)0.	4.017			попости	26, 110	•
Pb(Fe., Tao, 8)O3	4.011				26	
Pb(Ino.gNbo.s)Os	4.11				111	
Pb(Hoo.sNbo.s)Os	4.160	-	4.106	monoclinic	111	•
Fb(Lu _{0.8} Nb _{0.8})O ₃	4.152		4.098	monoclinic	111	
Fb(Lu _{0.8} Ta _{0.8})O ₃	4.153		4.107	monoclinic	111	
FD(Sco. 5 N Do. 5) O3	4.078		4.083	tetragonal	26, 112	
FD(50, 518, 5)U3	4.072				26, 112	
T D(T D0.51N D0.5) OB	4.10				96, 111,	
Pb(Yb, .Ta, .10.	4.13				112	
Sr(Co. Nb. 1)0.	3.93				94	
Sr(Coo. Sbo. s)Os	7.88	•		(NH4)3FeF	i	
<u>,</u>	0			structure	94	-
O1 (010.814100.8) U3	70.7	•		structure	88	
Sr(Cr _{0.8} Nb _{0.8})O ₃	3.9421		•		94, 107	

TABLE 2.2 (cont.)

A*+(B*+B*+)O3 (cont.)

Sr(Cr _{0.6} O8 _{0.6})O ₃ 7.84 5 (A) c (A) Remarks R			(m _{0.5} = 0.5) 3	3	(::::::::::::::::::::::::::::::::::::::	
7.84 (NH4,) ₃ FeF ₆ structure 7.82 (NH4,) ₃ FeF ₆ structure 7.862 (NH4,) ₃ FeF ₆ structure 7.82 (NH4,) ₃ FeF ₆ structure 3.94 (NH4,) ₃ FeF ₆ structure 7.843 (NH4,) ₃ FeF ₆ structure 7.843 (NH4,) ₃ FeF ₆ structure 7.843 (NH4,) ₃ FeF ₆ structure 8.05 (NH4,) ₃ FeF ₆ structure 8.071 structure (NH4,) ₃ FeF ₆ structure 8.27 structure (NH4,) ₃ FeF ₆ structure	Compound	a (Å)	(A) ó	c (Å)	Remarks	Refer- ences
### 17.82 NH4,13F6F,	Sr(Cr, Os, 1)O,	7.84			(NH,),FeF.	
7.82 (NH ₄) ₈ FeF ₆ structure 7.862 (NH ₄) ₈ FeF ₆ 8tructure 3.94 7.82 (NH ₄) ₃ FeF ₆ 8tructure 3.96 (NH ₄) ₃ FeF ₆ 8tructure 3.96 (NH ₄) ₃ FeF ₆ 8tructure 7.843 (NH ₄) ₃ FeF ₆ 8tructure 7.843 (NH ₄) ₃ FeF ₆ 8tructure 7.844 7.91 tetragonal 8.05 (NH ₄) ₃ FeF ₆ 8tructure 8.071 (NH ₄) ₃ FeF ₆ 8tructure 7.98 (NH ₄) ₃ FeF ₆ 8tructure		ļ			structure	18
7.862 (NH ₄) ₂ FeF ₆ 3.94 7.82 (NH ₄) ₂ FeF ₆ 8tructure 7.89 (NH ₄) ₂ FeF ₆ 8tructure 3.96 3.96 3.981 tetragonal 3.946 (NH ₄) ₃ FeF ₆ 8tructure 7.843 tetragonal 7.84 (NH ₄) ₃ FeF ₆ 8tructure 8.05 (NH ₄) ₃ FeF ₆ 8tructure 8.071 tetragonal 8.27 (NH ₄) ₃ FeF ₆ 8tructure 8.33 structure (NH ₄) ₃ FeF ₆ 8tructure 7.98 (NH ₄) ₃ FeF ₆ 8tructure	$\mathrm{Sr}(\mathrm{Cr_{0.8}Re_{0.8}})\mathrm{O_3}$	7.82			$(NH_4)_3FeF_8$	•
3.94 7.82 8.194 7.89 7.89 7.916 7.916 8.197 7.916 8.198 7.91 8.006 8.006 8.007 8.007 8.33 8.007 8.27 8.27 8.27 8.27 8.27 8.27 8.27 8.2	Sr(CrSb)O.	7.862	-		Structure (NH.) FAF.	87
3.94 7.82 (NH ₄) ₃ FeF ₆ structure 3.97 7.916 3.96 3.981 7.82 structure 3.946 7.82 structure 7.843 7.843 (NH ₄) ₃ FeF ₆ structure 7.843 7.91 4.0569 8.06 8.06 8.07 8.07 8.07 8.07 8.07 8.27 8.27 8.27 8.27 8.27 8.27 8.27 8.2	8) (9:0-)-0:0				structure	94, 108
7.89 7.89 (NH4,),FeF, structure 3.97 7.916 3.981 7.82 3.981 7.84 7.82 8.04 7.84 7.91 4.0569 8.06 8.07 8.07 8.07 8.33 8.07 8.27 8.27 8.27 8.27 8.27 8.27 8.27 8.2	$Sr(Cr_0.sTa_0.s)O_s$ $Sr(Cr_1.sV_1.s)O_s$	3.94			(NH.) FeF	19
7.89 (NH4,),FeF, 3.97 (NH4,),FeF, 4.916 structure 3.946 structure 3.946 (NH4,),FeF, 5.82 structure 7.84 (NH4,),FeF, 8.071 structure 8.071 structure (NH4,),FeF, 8.13 structure (NH4,),FeF, 8.27 structure	80/9.01.0.0	2			structure	58
7.89 (NH4,),FeF, 3.97 (NH4,),FeF, 5.916 structure 3.96 3.981 tetragonal 3.946 (NH4,),FeF, 5.82 structure 7.843 (NH4,),FeF, 8.06 structure 8.071 (NH4,),FeF, 8.33 structure (NH4,),FeF, 8.33 structure (NH4,),FeF, 8.34 (NH4,),FeF, 8.35 structure (NH4,),FeF, 8.37 (NH4,),FeF, 8.38 structure (NH4,),FeF, 8.39 (NH4,),FeF, 8.27 structure (NH4,),FeF, 8.39 structure (NH4,),FeF, 8.27 structure (NH4,),FeF, 8.27 structure (NH4,),FeF, 8.27 structure	Sr(Dy0.8 Ta0.6)O3					93
7.89 (NH4)3FeF, structure 3.97 (NH4)3FeF, structure 3.96 3.981 tetragonal 3.946 (NH4)3FeF, structure 7.843 (NH4)3FeF, structure 7.84 (NH4)3FeF, structure 8.07 (NH4)3FeF, structure 8.07 (NH4)3FeF, structure 8.27 (NH4)3FeF, structure	Sr(Er., Ta., S)O.					e e
3.97 7.916 (NH4,),FeF, structure 3.96 3.981 7.82 8.046 (NH4,),FeF, structure 7.843 (NH4,),FeF, structure 7.84 7.91 4.0569 8.06 8.07 8.06 8.07 8.07 8.07 8.10 8.10 8.27 8.27 8.27 8.27 8.27 8.27 8.27 8.27	Sr(Fe _{0.8} Mo _{0.8})O ₃	7.89			(NH,)3FeF	<u>.</u>
3.96 7.916 7.916 8.046 3.981 8.046 7.82 8.046 7.843 8.046 7.843 8.046 8.044,3-FeF ₆ 8.071 8.071 8.071 8.27 8.27 8.27 8.27 8.27 8.27 8.27 8.27		(structure	58
3.96 3.981 tetragonal 3.946 7.82 (NH4,)\$FeF6 structure 7.84 7.91 tetragonal 4.0569 8.06 structure 8.071 (NH4,)\$FeF6 structure 8.33 structure (NH4,)\$FeF6 structure 8.27 (NH4,)\$FeF6 structure 7.98 (NH4,)\$FeF6 structure (NH4,)\$FeF6 structure (NH4,)\$FeF6 structure (NH4,)\$FeF6 structure (NH4,)\$FeF6 structure (NH4,)\$FeF6 structure (NH4,)\$FeF6	Sr(Fe, Sb, k)O.	3.97 7.916		-	(NH,),FeF,	07
3.96 3.981 tetragonal 3.946 7.82 (NH4) ₃ FeF ₆ structure 7.84 7.91 tetragonal 4.0569 8.06 8.07 8.07 8.07 8.33 8.27 8.27 (NH4) ₃ FeF ₆ structure 8.33 8.27 (NH4) ₃ FeF ₆ structure 8.33 8.27 (NH4) ₃ FeF ₆ structure 7.98 (NH4) ₃ FeF ₆ structure 8.27 (NH4) ₃ FeF ₆ structure 8.27 (NH4) ₃ FeF ₆ structure 7.98 (NH4) ₃ FeF ₆ structure	9 (80)				structure	94, 108
3.946 7.82 structure 7.843 (NH4,)3FeF, structure 7.84 7.91 tetragonal 4.0569 (NH4,)3FeF, structure 8.071 structure (NH4,)3FeF, structure 8.33 structure (NH4,)3FeF, structure (NH4,)3FeF, structure 7.98 (NH4,)3FeF, structure (NH4,)3FeF, structure (NH4,)3FeF, structure (NH4,)3FeF, structure 7.98 (NH4,)3FeF,	Sr(Fe, 120, 8)08	3.96		3.981	tetragonal	113
4.0569 8.06 8.071 8.33 8.071 8.27 8.27 8.27 8.27 8.28 8.39 8.27 8.29 8.29 8.29 8.29 8.29 8.29 8.20 8.39 8.20 8.39 8.20 8.30 8.20 8.30 8.30 8.30 8.30 8.30 8.30 8.30 8.3	Sr(Geo.gN Do.s)O.	3.946			HAT (HN)	∄
7.843 (NH4,)3FeF, structure 7.84 7.91 tetragonal 4.0569 (NH4,)3FeF, 8.06 structure (NH4,)3FeF, structure 8.33 structure (NH4,)3FeF, structure 7.98 (NH4,)3FeF, structure 7.98 (NH4,)3FeF,	S (C C C C C C C S) C S	30.			structure	18
4.0569 4.0569 8.06 8.071 8.33 8.27 (NH4,)3FeF, structure	Sr(Ga _{0.8} Re _{0.8})O ₃	7.843			(NH,)3FeF,	·
4.0569 8.06 8.071 8.071 8.33 8.27 8.27 (NH4,)3FeF, 8tructure	} { {			,	structure	18
4.0569 8.06 8.071 8.071 (NH4,) ₃ FeF ₆ 8tructure 7.98 (NH4,) ₃ FeF ₆ 8tructure tetragonal	Sr(Gao. gSbo. s)Os	7.84		7.91	tetragonal	808
4.0569 8.06 8.071 8.071 (NH4) ₃ FeF ₆ 8tructure (NH ₄) ₃ FeF ₆ 8tructure 7.98 (NH ₄) ₃ FeF ₆ 8tructure tetragonal	Sr(Ho. Ta)O					2 6
8.06 8.071 8.071 8.33 8.27 8.27 (NH ₄) ₂ FeF ₆ 8tructure tetragonal	Sr(In, Nb, 1)0,	4.0569				107
8.071 (NH ₄) ₃ FeF ₆ 8.33 (NH ₄) ₃ FeF ₆ 8.27 (NH ₄) ₃ FeF ₆	Sr(Ino. 5 Oso. 5) Os	8.06			$(NH_4)_3$ FeF	
8.33 (NH _{4,3} FeF ₆ 8.23 (NH _{4,3} FeF ₆ 8.27 (NH _{4,3} FeF ₆ 8.27 (NH _{4,3} FeF ₆ 8tructure 7.98 (NH _{4,3} FeF ₆ 8tructure	ر. ط در الم	i c			structure	18
8.33 (NH ₄) ₃ FeF ₆ structure (NH ₄) ₃ FeF ₆ structure (NH ₄) ₃ FeF ₆ structure tetragonal	OI(1110.5 TO 9.5) U3	8.071			Structure	18
8.27 (NH ₄) ₃ FeF ₆ structure (NH ₄) ₃ FeF ₆ structure structure tetragonal	$Sr(In_0.sU_0.s)O_3$	8.33			(NH4)3FeF)
8.27 (NH4,)3FeF, 8tructure 7.98 (NH4,)3FeF, 8tructure tetragonal		· · · · · · · · · · · · · · · · · · ·			structure	104
7.98 (NH ₄) ₃ FeF ₆ structure tetragonal	Sr(La _{0.8} Ta _{0.8})O ₃	8.27			(NH4)3FeF	6
7.98 (NH ₄) ₃ FeF ₆ structure tetragonal	Sr(L,, To)O				structure	# 65
structure tetragonal	Sr(Mn, Mo, 1)0.	7.98	·		(NH,),FeF,	3
tetragonal					structure	28
tetragonal	Sr(Mn0.8Sb0.8)O3	,		-		114
10110801101	Sr(Nd _{0.8} 1'8 _{0.8})O ₃ Sr/Ni Sh \O					28
	OL(1/10.6/3/0.8/)O3					K

PREPARATION OF PEROVSKITE-TYPE COMPOUNDS

(cont.)	
A*+(B _{0.5} + B _{0.5})O ₃	
(cont.)	
LABLE 2.2 (

Compound	a (A) b (A) c (A)	b (A)	c (A)	Remarks	Refer- ences
Sr(Rho., Sbo.,)O3	5.77	6.55	6.55 7.99	orthorhombic	84
SI (SC0.8 OS0.8) O3	8.0z			(NH4)3F6F6 structure	18
Sr(Sc _{0.8} Re _{0.8})O ₃	8.02			(NH4),FeF.	
O"(0, Th. 0, 1)				structure	× ×
Sr(Tm _{0.8} Ta _{0.8})O ₃					88
Sr(Yb _{0.8} Ta _{0.8})O ₃					83

$A^{2+}(B_{0.5}^{2+}B_{0.5}^{6+})O_3$

Ba(Ba, Og 10.	8		76.0	+0+	91
Ba(Ba _{0.8} Re _{0.8})O ₃	8.65		8.33	tetragonal	18
Ba(Ba _{0.5} U _{0.5})O ₃	8.83			(NH,),FeF,	
				structure	17
Ba(Ba _{0.8} W _{0.8})O ₃	8.6			(NH,)3FeF	
				structure	16
Ba(Ca,, Mo,, s)O3	8.355			(NH4),FeF	
				structure	16
Ba(Ca _{0.8} Os _{0.8})O ₃	8.362			(NH,)3FeF,	
				structure	18
Ba(Ca _{0.8} Re _{0.8})O ₂	8.356			(NH,),FeF,	
				structure	18, 115
Ba(Ca., Te,., O3	8.393			(NH,)3FeF	
				structure	108
Ba(Ca _{0.8} U _{0.8})O ₃	8.67			(NH,)3FeF	
				structure	17
Ba(Ca _{0.8} W _{0.5})O ₃	8.39			(NH,)3FeF	
				structure	15, 16
Ba(Cdo.,OSo.,s)O3	8.326			(NH,),FeF	
				structure	18
Ba(Cd _{0.5} Re _{0.5})O ₃	8.322			(NH,),FeF,	
				structure	18, 115
Ba(Cd _{0.8} U _{0.6})O ₃	6.13	8.64	6.07	orthorhombic	17
Ba(Co,,,Mo,,,)O,	4.0429				107
Ba(Co,, Re,, s)O,	8.086			(NH,),FeF,	
,				structure	18, 115
Ba(Co,, U,, s)O3	8.374			(NH,),FeF	
				structure	17
		-	1		

STRUCTURE OF PEROVSKITE-TYPE COMPOUNDS TABLE 2.2 (co

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V 2+/D	

Ba(Co _{0.5} W _{0.5})O ₃ Ba(Co _{0.5} W _{0.5})O ₃ Ba(Co _{0.5} U _{0.5} O ₃) Ba(Co _{0.5} U _{0.5} O ₃) Ba(Fe _{0.5} U _{0.5} O ₃) Ba(Mg _{0.5} Re _{0.5})O ₃ Ba(Mg _{0.5} U _{0.5} O ₃) Ba(Ni _{0.5} Me _{0.5} O ₃) Ba(Ni _{0.5} U _{0.5} O ₃) Ba(Ni _{0.5} U _{0.5} O ₃) Ba(Ni _{0.5} U _{0.5} O ₃) Ba(Si _{0.5} U _{0.5} O ₃ O ₃) Ba(Si _{0.5} U _{0.5} O ₃ O ₃) Ba(Si _{0.5} U _{0.5} O ₃ O ₃) Ba(Si _{0.5} U _{0.5} O ₃ O ₃) Ba(Si _{0.5} U _{0.5} O ₃ O ₃) Ba(Si _{0.5} U _{0.5} O ₃ O ₃ O ₃ O ₃) Ba(Si _{0.5} U _{0.5} O ₃	Compound	a (Å)	b (A)	c (Å)	Remarks	References
8.297 (NH ₄) ₈ FeF ₆ 8.18 (NH ₄) ₈ FeF ₆ 8.112 (NH ₄) ₈ FeF ₆ 8.113 (NH ₄) ₈ FeF ₆ 8.082 (NH ₄) ₈ FeF ₆ 8.082 (NH ₄) ₈ FeF ₆ 8.082 (NH ₄) ₈ FeF ₆ 8.083 (NH ₄) ₈ FeF ₆ 8.084 (NH ₄) ₈ FeF ₆ 8.085 (NH ₄) ₈ FeF ₆ 8.086 (NH ₄) ₈ FeF ₆ 8.089 (NH ₄) ₈ FeF ₆ 8.099 (NH ₄) ₈ FeF ₆ 8.099 (NH ₄) ₈ FeF ₆ 8.099 (NH ₄) ₈ FeF ₆ 8.096 (NH ₄) ₈ FeF ₆ 8.009 (NH ₄) ₈ FeF ₆	Ba(Co., Wo., s)Os	8:098			(NH4) ₈ FeFe	15, 107
8.18 8.84 tetragonal 8.05 8.312 8.312 (NH ₄) ₂ FeF ₆ 8.133 8.133 (NH ₄) ₂ FeF ₆ 8.133 (NH ₄) ₂ FeF ₆ 8.134 (NH ₄) ₂ FeF ₆ 8.138 (NH ₄) ₂ FeF ₆ 8.138 (NH ₄) ₂ FeF ₆ 8.138 (NH ₄) ₃ FeF ₆ 8.149 (NH ₄) ₃ FeF ₆ 8.140 (NH ₄) ₄ FeF ₆ 8.140 (Ba(Cro., Uo., s)O3	8.297		•	structure (NH4)3FeF6	17
8.18 8.05 8.05 8.10 8.112 8.12 8.133 8.133 8.133 8.133 8.14,1,3,FeF, 8.13 8.13 8.14,1,3,FeF, 8.14,1,3,FeF, 8.18 8.18 8.18 8.18 8.18 8.18 8.18 8.1	, ,				structure	;
8.132 (NH4,),FeF, 8.133 (NH4,),FeF, 8.108 (NH4,),FeF, 8.108 (NH4,),FeF, 8.13 (NH4,),FeF, 8.13 (NH4,),FeF, 8.13 (NH4,),FeF, 8.13 (NH4,),FeF, 8.14 (NH4,),FeF, 8.18 (NH4,),FeF, 8.28 (NH4,),FeF, 8.29 (NH4,),FeF, 8.	Ba(Fe,, Re,, s)O3 Ba(Fe,, Re,, s)O3	8.18 8.05		8.8 4	tetragonal (NH,) ₃ FeF,	17
8.133 (NH4,),FeF, structure 8.08 (NH4,),FeF, structure 8.13 (NH4,),FeF, structure 8.13 (NH4,),FeF, structure 8.099 (NH4,),FeF, structure 8.09 (NH4,),FeF, structure 8.04 (NH4,),FeF, structure 8.36 (NH4,),FeF, structure 8.60 (NH4,),FeF, structure 8.84 (NH4,),FeF, structure 8.85 (NH4,),FeF, structure 8.095	Ba/Fe1110.	8.312			structure (NH.) FeF	18, 116
8.133 (NH4,13F9F, structure 8.08 (NH4,13F9F, structure 8.13 (NH4,13F9F, structure 8.13 (NH4,13F9F, structure 8.09 (NH4,13F9F, structure 8.04 (NH4,13F9F, structure 8.36 (NH4,13F9F, structure 8.60 (NH4,13F9F, structure 8.60 (NH4,13F9F, structure 8.60 (NH4,13F9F, structure 8.86 (NH4,13F9F, structure 8.86 (NH4,13F9F, structure 8.89 (NH4,13F9F, structure 8.096 (NH4,13F9F)	20 (2 (0.8) (8				structure	17
8.08 8.082 8.082 8.082 8.13 8.13 8.13 8.13 8.099 8.18 8.18 8.09 8.18 8.18 8.04 8.18 8.04 8.18 8.04 8.18 8.04 8.18 8.36 8.36 8.40 8.40 8.40 8.40 8.40 8.40 8.40 8.40	Ba(Fe, 5Wo.5)O3	8.133		•	(NH,)3FeF,	1
8.082 structure structure s.13 structure s.13 structure s.381 structure s.089 structure s.089 structure s.089 structure s.08 s.04 structure s.08 s.04 structure s.08 s.08 structure s.08 s.09 structure structure s.09 structure structure s.09 structure structure structure s.09 structure s	Ba(Mgo.sOso.s)Os	8.08			(NH4),FeF,	01
8.13 structure 8.381 structure 8.099 structure 8.18 structure 8.18 structure 8.62 (NH4,) FeF, 8.04 structure 8.336 (NH4,) FeF, 8.336 structure 8.336 (NH4,) FeF, 8.43 structure 8.43 structure 8.43 structure 8.43 structure 8.44 structure 8.45 structure 8.48 structure 8.48 structure 8.49 structure 8.40 structure 8.50 (NH4,) FeF, 8 structure 8.60 (NH4,) FeF, 8 structure 8.60 (NH4,) FeF, 8 structure 8.095 structure 8.095 structure 8.095 structure 8.095 structure 8.095 structure 8.095 structure	Ba(Mg, Re,)O.	8.082			structure (NH,),FeF,	18
8.381 (NH4,)gFeF, structure 8.099 (NH4,)gFeF, structure 8.18 (NH4,)gFeF, structure 8.52 (NH4,)gFeF, structure 4.0225 (NH4,)gFeF, structure 8.336 (NH4,)gFeF, structure 8.336 (NH4,)gFeF, structure 8.43 8.72 tetragonal 8.60 (NH4,)gFeF, structure 8.543 8.72 tetragonal 8.60 (NH4,)gFeF, structure 8.72 tetragonal 8.60 (NH4,)gFeF, structure 8.60 (NH4,)gFeF, structure 8.095 (NH4,)gFeF, structure	O("E ")() B	6.			structure	18, 115
8.381 (NH ₄),FeF ₆ 8.099 (NH ₄),FeF ₆ 8.18 (NH ₄),FeF ₆ 8.18 (NH ₄),FeF ₆ 8.04 structure 8.336 (NH ₄),FeF ₆ 8.43 structure 8.43 structure 8.43 structure 8.43 structure 8.44 structure 8.56 (NH ₄),FeF ₆ 8.56 (NH ₄),FeF ₆ 8.57 tetragonal 8.60 structure 8.60 (NH ₄),FeF ₆ 8.70 tetragonal 8.70	D&(ME0.5 1 e0.6) U8	6.13			(IN TA 4) 3F GF g structure	108, 116
8.099 (NH4), FeF, 8.18 (NH4), FeF, 8.18 (NH4), FeF, 8.18 (NH4), FeF, 8.04 (NH4), FeF, 8.04 (NH4), FeF, 8.036 (NH4), FeF, 8.066 (NH4), FeF, 8.13 (NH4), FeF, 8.13 (NH4), FeF, 8.14 (NH4), FeF, 8.15 (NH4), FeF, 8.16 (NH4), FeF, 8.17 tetragonal 8.60 (NH4), FeF, 8.18 (NH4), FeF, 8.19 (NH4), FeF, 81095 (NH4), FeF, 81100000000000000000000000000000000000	Ba(Mgo.sUo.s)O3	8.381			(NH,)3FeF	
8.18 (NH ₄) ₂ FeF ₆ 8.52 structure 4.0225 8.04 structure 8.336 (NH ₄) ₃ FeF ₆ structure 8.336 structure 8.066 (NH ₄) ₃ FeF ₆ structure 8.43 structure 8.43 8.72 tetragonal 8.60 tetragonal 8.60 (NH ₄) ₃ FeF ₆ structure 8.50 tetragonal 8.60 structure 8.50 (NH ₄) ₃ FeF ₆ structure 8.60 (NH ₄) ₃ FeF ₆ structure 8.60 (NH ₄) ₃ FeF ₆ structure 8.60 (NH ₄) ₃ FeF ₆ structure 8.095 structure 8.095 structure	Ba(Mgo.s Wo.s)O3	8.099			Structure (NH4)3FeF,	11
8.52 (NH4,)3FeF, 8.04 (NH4,)3FeF, 8.04 (NH4,)3FeF, 8.036 (NH4,)3FeF, 8.066 (NH4,)3FeF, 8.43 (NH4,)3FeF, 8.60 (NH4,)3FeF, 8.72 tetragonal 8.60 (NH3,)3FeF, 8.72 tetragonal 8.60 (NH4,)3FeF, 8.73 (NH4,)3FeF, 8.74 tetragonal 8.60 (NH4,)3FeF, 8.75 tetragonal 8.60 (NH4,)3FeF, 8.76 tetragonal 8.77 tetragonal 8.78 tetragonal 8.84 structure 8.95 structure 8.095 structure	Ba(Mn. Ba. 10.	α			structure	16, 16
8.62 (NH ₄) ₂ FeF ₆ 8.04 (NH ₄) ₃ FeF ₆ 8.04 (NH ₄) ₃ FeF ₆ 8.336 (NH ₄) ₃ FeF ₆ 8.066 (NH ₄) ₃ FeF ₆ 8.43 8.72 tetragonal 8.60 8.29 tetragonal 8.84 (NH ₄) ₃ FeF ₆ 8.72 tetragonal 8.60 (NH ₄) ₃ FeF ₆ 8.72 tetragonal 8.60 (NH ₄) ₃ FeF ₆ 8.73 tetragonal 8.60 (NH ₄) ₃ FeF ₆ 8.74 tetragonal 8.75 tetragonal 8.60 (NH ₄) ₃ FeF ₆ 8.75 tetragonal 8.76 (NH ₄) ₃ FeF ₆ 8.77 tetragonal 8.78 (NH ₄) ₃ FeF ₆ 8.79 tetragonal 8.70 tetragonal	20 (21 TO 10 10 10 10 10 10 10 10 10 10 10 10 10	3			structure	18, 115
4.0225 8.04 8.04 8.136 8.336 8.336 8.086 8.086 8.43 8.43 8.60 8.43 8.60 8.84 8.60 8.84 8.60 8.84 8.60 8.72 tetragonal 8.86 8.72 tetragonal 8.86 8.72 tetragonal 8.86 8.72 tetragonal 8.86 8.73 8.72 tetragonal 8.86 8.73 8.74 8.75 tetragonal 8.86 8.75 tetragonal 8.86 8.76 8.77 8.78 8.79 8.79 8.79 8.70 8.79 8.70 8.70 8.70 8.70 8.70 8.70 8.70 8.70	Ba(Mno.5Uo.5)O3	8.52			(NH4)3FeFe	}
8.04 (NH4,)FeF6 8.336 (NH4,)FeF6 8.066 (NH4,)FeF6 8.43 8.72 tetragonal 8.60 8.29 tetragonal 8.84 (NH,)FeF6 8.096 (NH4,)FeF6 8.096 (NH4,)FeF6 8.095 structure	Ba(Ni, Mo, ,)O,	4.0225			structure	107
8.336 (NH _a) ₃ FeF _e structure 8.066 (NH _a) ₃ FeF _e structure 8.43 8.72 tetragonal 8.60 8.29 tetragonal 8.84 (NH _a) ₃ FeF _e structure 8.5 (NH _a) ₃ FeF _e structure (NH _a) ₃ FeF _e structure (NH _a) ₃ FeF _e structure 8.096 (NH _a) ₃ FeF _e structure	Ba(Ni _{0.5} Re _{0.5})O ₃	8.04			(NH,),FeF,	
8.066 (NH4,),FeF, structure 8.43 8.72 tetragonal 8.60 (NH,),FeF, structure 8.5 structure 8.5 structure 8.095 (NH4,),FeF, structure 8.095 structure	Ba(Ni _{0.8} U _{0.8})O ₈	8.336			structure (NH4),FeF	18, 116
8.43 8.72 tetragonal 8.60 8.29 tetragonal 8.84 (NH_0)_FEF_ 8.5 (NH_0)_FEF_ 8.5 (NH_0)_FEF_ 8.095 (NH_0)_FEF_ 8tructure (NH_0)_FEF_ 8tructure 8.095 (NH_0)_FEF_ 8tructure	Be(N; W)O	8 088			Structure	17
8.43 8.72 tetragonal 8.60 8.29 tetragonal 8.84 (NH ₄) ₃ FeF ₆ 8.5 (NH ₄) ₄ FeF ₆ 8.095 structure 8.095 structure	20(110.8 110.8)				structure	15, 107
8.60 8.29 tetragonal 8.84 (NE ₄) ₄ FeF ₆ structure 8.5 (NH ₄) ₄ FeF ₆ structure 8.095 (NH ₄) ₄ FeF ₆	Ba(Pb _{0.8} Mo _{0.8})O ₈ Ba(Sr. OS)O.	8.43		8.72	tetragonal	117
8.5 (NH _d) ₃ FeF ₆ 8.5 (NH _d) ₃ FeF ₆ 8.095 (NH _d) ₃ FeF ₆	Ba(Sr _{0.8} Re _{0.8})O ₈	8.60		8.29	tetragonal	18, 116
8.5 (NH ₄) ₃ FeF ₆ structure (NH ₄) ₃ FeF ₆ structure	Da(Sr _{0.5} ∪ _{0.5}) ∪ ₃	8.8 4			(IN H.4)3F 8.F.6 structure	17
8.095 (NH _d)3FeF _e structure	$Ba(Sr_{0.5}W_{0.5})O_s$	8.5			(NH4) ₈ FeF	91
structure	Ba(Zno. GOso. B)Os	8.095			Structure (NH4),FeF	2
					structure	18

Commonned	7 (1)	1, 1, 1,			Refer-
Compound	a (A)	(A)	c (A)	Remarks	ences
$\mathrm{Ba}(\mathrm{Zn}_{0.5}\mathrm{Re}_{0.5})\mathrm{O}_3$	8.106			(NH ₄) ₈ FeF ₈	
Ba(Zn, U, 10.	8.397			structure	18, 115
8 (0:0 - 0:0 -)				structure	17
$\mathrm{Ba}(\mathrm{Zn_{0.5}W_{0.5}})\mathrm{O_3}$	8.116			(NH4)3FeFs	
Ca(Ca. Os)O.	5 73	1	o u	structure	15
Ca(Ca, Re, O	5.67	8.05	7.00	orthorhombie	20
Ca(Ca _{0.8} W _{0.8})O ₃	8.0	}	?	(NH4),FeF	07
4				structure	16
Ca(Cdo.s.Keo.s)O3	5.64	7.99	5.73	orthorhombic	18
Ca(Co, a CBo, s) Ca	0.47	7.70	5.59	orthorhombic	18
Ca(Co).5 Dej. 5) C3	5.40	7.71	5.58	orthorhombic	18
Ca(Mo. Re. 10)	0.41	7.09	50.0	orthorhombic	18, 115
Ca(Mg, W,)O	7.7	:	0.00	Orthornomole (NH) FeF	2 0
				Structure Structure	9
$Ca(Mn_0, Re_0, s)O_s$	5.52	7.82	5.55	orthorhombic	2 0
Ca(Nio., Reo., s)O,	5.45	7.67	5.65	orthorhombic	18
$Ca(Sr_0{\delta}W_{0.\delta})O_3$	8.1			(NH4)3FeF6	
Dh/Ce III 10				structure	16
Pb(Cd. W. 10.	4 150	חאר אוסר א	150	0000 T 0	96
Pb(Co., W.,)O.	20	101.1	201.		011
Pb(Mg,,Te,,)O,	7.99			(NH.), FAF.	611
			•	structure	116
Pb(Mgo.s Wo.s)O3	4.0				95, 96
Sr(Ca _{0.8} Mo _{0.5})O ₃					117
Sr(Ca _{0.5} Os _{0.5})O ₃	8.21			(NH,)3FeF	•
Sr(Ca. Re 10	7. A.	60		structure	2 2
Sr(Ca, U.)O.	6.06		5 93	orthorhombie	7 F
Sr(Ca0.5W0.5)O3	8.2		}	(NH,),FeF,	-
		_		structure	16
Sr(Cd, , Re, ,)O3	5.73		5.81	orthorhombic	18
Sr(Ca _{0.8} C _{0.8})O ₃	6.03	8.42	5.91	orthorhombic	17
Sr(Co.5-100.5) C3	1000.0	3 E	4016.6	1-1-1-1	107, 113
Sr(Co. Re. 10.			78.7	tetragonal	× 2
Sr(Co, 5U, 5)O3	8.19			(NH,), FeF.	2
					;

TABLE 2.2 (cont.)		A2+(B2+B6+)O3 (cont.))O3 (cont.)	
Compound	a (Å)	b (Å)	c (Å)	Remarks	Refer- ences
$Sr(Co_{o,s}W_{o,s})O_{s}$ $Sr(Cr_{o,s}U_{o,s})O_{s}$	7.89		7.98	tetragonal (NH ₄) ₈ FeF ₄	16, 113
; ;				structure	17
$\mathrm{Sr}(\mathrm{Cu}_0{\mathfrak{s}}\mathrm{W}_{\mathfrak{o}{\mathfrak{s}}})\mathrm{O}_{\mathfrak{s}} \ \mathrm{Sr}(\mathrm{Fe}_{\mathfrak{o}{\mathfrak{s}}}\mathrm{Os}_{\mathfrak{o}{\mathfrak{s}}})\mathrm{O}_{\mathfrak{s}}$	7.66		8.40	tetragonal (NH4)3FeF	94
£	-		,	w	18
$\mathrm{Sr}(\mathrm{Fe_{0.5}Ke_{0.5}})\mathrm{O_{3}}$ $\mathrm{Sr}(\mathrm{Fe_{0.5}U_{0.5}})\mathrm{O_{3}}$	8.11		7.89	tetragonal (NH4)3FeF	18, 116
$Sr(Fe_{0.8}W_{0.8})O_s$	7.96			structure	17
				structure	94
$Sr(Mg_0,sMo_0,s)O_3$	7 88		90	1000000	117
Sr(Mg0.8Re0.8)O3	7.88		7.94	tetragonal	9 8
Sr(Mgo., Teo., 5)O3	7.84			(NH4)3FeF	<u>}</u>
				structure	116
Sr(Mgo.sUo.s)Os	8.19			(NH,)3FeFg	
Sr(Ma. W.)O.	10			structure	17
0.6/41-50.5 11 0.5/03	9:			(IN IL 4)3F OF 8	91
$\mathrm{Sr}(\mathrm{Mn_{0.8}Re_{0.8}})\mathrm{O_3}$	8.01			(NH,)3FeF	?
77 TL	6			structure	18
Or(14th _{0.5} U _{0.5}) U ₃	87.8			(NH ₄) ₃ FeF ₆	
$Sr(Mn_{0.6}W_{0.6})O_3$	8.01			(NH,),FeF.	7
				structure	94
Sr(Ni _{0.5} Mo _{0.5})O ₃	3.9237		3.9474	tetragonal	107, 113
Sr(N10.5 Reo.5) O.	7.85		7.92	tetragonal	18
Of (21.0.8 0.8) (3	91.0			(INTA)3FEFF	,
Sr(Ni, W. 10.	7 86		107	structure ************************************	17
0.6/4.40.5 11 0.6/ 0.8	5		16.1	rentagonai	113, 107,
Sr(Pbo.sMoo.s)Os					117
$Sr(Sr_0.sOs_0.s)O_3$	8.32		8.12	tetragonal	18
Sr(Sr _{0.8} Re _{0.8})O ₃	8.41		8.13	tetragonal	18
Sr(Sr ₀ , s U ₀ , s)O ₃ Sr(Sr ₀ , W ₀ , 1)O ₂	6.22 8.22	8.65	6.01	orthorhombic	17
0.0 1 0.0 V 3	•			structure	16

tetragonal tetragonal orthorhombic (NH₄)₃FeF₆ structure tetragonal tetragonal

8.01

7.89

Sr(Zn_{0.8}Re_{0.8})O₃ Sr(Zn_{0.8}W_{0.8})O₃

 $A^{2+}(B_{0.5}^{1+}B_{0.5}^{7+})O_{3}$

TABLE 2.2 (cont.)

38

TABLE 2.2 (cont.)

 $A^{\sharp+}(B_{0.28}^{1+}B_{0.75}^{6+})O_3$

Refer-ences

Remarks

b (A) | c (A)

a (Å)

Compound

4.137

Ba(Na_{0.28}Ta_{0.78})O₂ Sr(Na_{0.28}Ta_{0.78})O₈

27

			•	~-	•		•					•			1	щ		щ		Ø			F
Refer- ences		108		18	91	0	18, 108		18		18		18		18		18		18		18		18
Remarks	(NH4)3FeF	structure	(NHA)3FeF's	Structure (NH.), FeF	Structure	(NH4)3FeF	structure	PARTE (FOR)	structure	(NH4)3FeF	structure	(NH,)3FeF,	structure	(NH,)3FeF	structure	(NH,)3FeF,	structure	(NH,)3FeF	structure	(NH4)3FeF	structure	(NH,),FeF.	structure
c (A)																							_
b (Å)																							
a(A) $b(A)$ $c(A)$	8.46	8.100	201:0	8.118		8.33	8.291	1	906 8	0.53.0	c c	7.03	0	.00	t	98.7	i c	1.8.7	,	8.13	,	8.13	
Compound	$\mathrm{Ba}(\mathrm{Ag_{0.8}I_{0.8}})\mathrm{O_{a}}$	Ba(Li, Og, 1)0,	80/0:0-0:0	$\operatorname{Ba}(\operatorname{Li}_{\mathfrak{d.g}}\operatorname{Re}_{\mathfrak{d.g}})\operatorname{O}_{\mathfrak{g}}$; ;	$Ba(Na_{\mathfrak{d},\mathfrak{s}^{L_{\mathfrak{g}},\mathfrak{s}}})O_{\mathfrak{s}}$	Ba(Nao.sOsos)O,		Ba(Na., Re., 10	0.8-1-0.8/	Ca(T.i. On 10	0.80.00.8	Ca(T,i. Be 10	0.8+20.8/03	Sr(T.i. O. vo	0.80.80.80.80.80.80.80.80.80.80.80.80.80	Sr(Li Bo 10	80/8-00.8-00.8/Og	Sr(Ne O. O.	01 (100.6 O80.8) U3	Sr(No Do	Of (1100.8100.8) O3	

104

(NH4)₈FeF

8.551

Ba(Ino., Uo., O2.78

A(B_{0.5} B_{0.5})O_{2.76}

structure

2.3. MADELUNG ENERGY

The calculation of the binding energy of a crystal is one of the fundamental problems in the theory of solids. In this calculation, the basic assumption in the theory of cohesive energy of ionic crystals is that the solid can be considered as a system of positive and negative ions. In the NaCl strucetc. The Coulomb energy of the sodium ion in the field of sodium ion is surrounded by 6 Cl⁻ ions at a distance L, 12 Na⁺ ions at a distance $L\sqrt{2}$, 8 Cl⁻ ions at a distance $L\sqrt{3}$, ture, the shortest interionic distance is given by L. Each all other ions is therefore,

$$U_M = -\frac{e^2}{L} \left(\frac{6}{\sqrt{1}} - \frac{12}{\sqrt{2}} + \frac{8}{\sqrt{3}} - \frac{6}{\sqrt{4}} + \frac{24}{\sqrt{5}} \right)$$

 $A^{3+}(B_{2}^{2}+B_{4}^{4}+)0$

	94	94	19	93	26,94	26	19, 96		26	26	26 94	96	2 -	n 0	270	19
	orthorhombic	monoclinic		.•				(NH4)3FeF	structure				•	*.		
0.3 -0.6 / 3																_
;	·	-	0.50	78.7	5	18.7	3.932	7.86		7.84	7.90	7.90	3.93	7.97	000	0.50
	La(Co., Ir.,)O. La(Cir., Ir.,)O.	La (Mg Co)	T. M. T. 10	T.8 (Mg. NT.)	La(Mg B 1)	To (Ma Tr.)	To (Mr. T-)	La(Mn0.8170.8)U3	To (Mr. D.,)	Talmio.grup.g) Us	LE(1N10.6IF0.6)O3	La(N10.8 Kuo.8)O3	La(Nio. 8Tio. 8)O3	La(Zno.,Ruo.,)O,	Nd/Me. Ti. 10	60.90.8/03

40

where e is the charge per ion. The coefficient of e^2/L is a pure number, determined only by the crystal structure and called the Madelung constant, M_L .

The Madelung constant for the cubic perovskite structure is 24.755. Using the equation

$$U_M = -M_L e^2/L$$

the Madelung energy for an assumed ideal structure for CaTiO, calculates out to be 4280.5.(117)

delung energy of idealized ordered cubic perovskites. These plus $n\pm$ charges. The ordered structure for these compounds calculations were based on formulas A2+B4-"B4+"O6 which is twice the A(B',B',')O₃ type formulas used throughout this report. In their formula $4\pm n$ denoted a net charge of 4^+ Rosenstein and Schor(121) determined the superlattice Mais shown in Fig. 2.5.

The method used for computing the change in Madelung energy due to charge ordering of the B ions is in accordance with the technique of superposition employed by Templeton, $^{(122)}$ since the ordering charges n^+ and n^- for an ordered Using $M_L=1.74756$ and $L=\frac{1}{4}$ supercell edge they derived perovskite were at alternate sites in a rocksalt structure. the Madelung energy due to charge ordering,

$$U_M = -1.74756 \times 331.984 \times 0.5 \, n^2/L.$$

The Madelung energies are given in Table 2.3 below.

TABLE 2.3. Table of Madelung Energies for Idealized Ordered Perovskite Compounds (after Rosenstein and Schor(121))

Kcal mole-1 total	8661 8199 8755 9383
Madelung energy ordering	142 577 1286
Supercell edge (Å)	7.68 8.16 8.04 8.12
Compound	Ca ₂ Ti ₂ O ₆ Ba ₂ Sc ³⁺ Re ⁶⁺ O ₆ Ba ₂ Ni ²⁺ Re ⁶⁺ O ₆ Ba ₂ Li ¹⁺ Re ⁷⁺ O ₆
ε	32 3

The results showed that the energy due to charge ordering was in some cases a significant fraction of the total Madelung

energy especially where the charge difference in the B ions

was large.

from 0.90 to 1.10. Least-squares fits expressing the Madelung constant as a function of $(1-\alpha)$ also were reported. The results gonal perovskite structures for axial ratios $\alpha = c/a$ varying In a later study, Saltzman and Schor (123) calculated the Madelung energy of tetragonal perovskite structure. Madelung constants were determined for the 4-4, 3-5, and 1-7 tetraare given in Table 2.4.

TABLE 2.4. Table of Madelung Energies (after Salzman and Schor(123)

Total Madelung energy	8236 8244 8284 8289 8280 8896 8946 8934 8911 8631 8631 8638
Madelung energy ordering	543 544 544 544 545 580 580 580 560 580
Madelung constant	53.70 53.70 52.43 52.78 52.94 52.87 52.87 53.65 53.65
$\alpha = c/a$	0.963 0.963 0.964 1.034 1.004 1.008 1.008 0.976 0.967
Compound	Ba,BaOsO, Ba,BaReO, Ba,SrReO, Ba,SrOsO, Sr,CoReO, Sr,MgOsO, Sr,MiReO, Sr,NiReO, Sr,Sr,NiReO, Sr,Sr,ReOo, Sr,Sr,ReOo, Sr,SrReOo, Sr,SrReOo,

2.4. Ionio Radii

appear to differ considerably in radius in the structure of perovskite-type compounds as compared with those of Ahrens. In addition, the ionic radii of W5+ and O87+ were The ionic radii of the ions as given by Ahrens and as calcu-The ionic radii of the B ion was obtained in ABO₃-type compounds, while those of B' and B" were obtained in complex perovskite compounds. There are a number of ions which lated in perovskite-type compounds are listed in Table 2.5. calculated for the first time.

TABLE 2.5. Ionic Radii

TABLE 2.5 (cont.)

	ı					
	B.'§		·			-
tonic naari	B′‡		0.99 0.97 0.73	0.74 0.80 0.69 1.20	1.12	
5	Bţ	· · · · · · · · · · · · · · · · · · ·				0.558
ander.	Abrens	1.26 1.37 1.67 0.96 1.80 0.62 1.33 0.68 0.94	0.89 1.34 0.35 0.99 0.97 0.73 0.73	0.73 0.67 0.80 0.69 1.20 0.80	1.43 0.93 1.12 0.76 0.95 0.74	1.18 0.65 1.07 0.58
		Ag1+ Ca Ca Cu Cu Fr Fr I I	Agg*+ Bag Coa	Hes Mas Mas No. Pd Pd	Ra Sr Ti Zn	Ac³+ Al Am As

B''\$								
B'‡	1.06	0.94 0.91 0.99 0.63	0.93	1.14 0.86 0.67 1.04	1.02	0.74	0.90	88.0
m m	0.56	0.628	0.714	0.625		0.686	0.625	
Ahrens	0.85 0.23 0.93 1.07 0.63	0.92 0.98 0.08 0.64	0.97 0.91 0.81	0.85 0.66 1.10	1.13 1.06 1.06 1.08 0.68	0.81 1.00 0.93 0.76	0.95 0.87 0.74 0.92	0.86 0.92 0.16
	් ර්රීර්කික ^{ති}	Er Fe Ga	r n gq	Lu N N N N O O O	7 P.	S S m T I	E E A	Yb Am ⁴⁺ C

TABLE 2.5 (cont.)

TABLE 2.5 (cont.)

B''8		
B/‡		0.69
B‡		
Abrens	0.94 0.63 0.68 0.69 0.09 0.98 0.98 0.92 0.93 0.65 0.65 0.67 0.71 0.70 0.70 0.63 0.63	0.46 0.74 0.69 0.35 0.89
	Z≪≺diijigssssarrrr R≪≺diijigsssssarrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrr	Ass*+ Nb Pa Pa Sb

B′′§ $0.68 \\ 0.59 \\ 0.66$ 0.62 0.65 B,‡ Ħ Abrens $0.68 \\ 0.69$ 0.62 0.62 0.67 0.52 0.30 0.42 0.65 0.80 $0.62 \\ 0.39$ $0.27 \\ 0.08 \\ 0.50$ 0.46 Atr Br Ci

† Calculated for ABO₃-type compounds by S. Geller, Acta Cryst. 10, 248 (1957).
‡ Ionic radii of B³⁺: Calculated from complex perovskite compounds A(B³;Tra_{0.3})O₃ by F. Galasso et al., UACRL D910269-5, Final Report, July 1965. Ionic radii of B³⁺: Calculated from complex perovskite compounds A(B_{0.33} Tra_{0.67})O₃₂₃.
§ Calculated from A(B_{0.5}B_{0.5})O₃ perovskite.

type compounds.

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CHAPTER 5

FERROELECTRICITY

ed on titanates by von Hippel et al. at MIT under wartime potential is applied to a capacitor containing a ferroelectric restrictions.(1) After the war, the work was released along used because these materials are analogous in some ways to tential, or polarization and field, produces a hysteresis loop on a cathode-ray oscilloscope (see Fig. 5.1). Ferromagnetic with studies conducted in England, (2-4) Russia (5-7) and Japan. (8) As a result of these investigations, barium titanate was found to be a ferroelectric.(?) The term ferroelectric was ferromagnetic materials. For example, when an alternating material, the instantaneous relation between charge and po-During World War II extensive investigations were conduct-

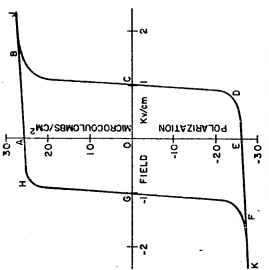


Fig. 5.1. Ferroelectric hysteresis loop.

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 $\tilde{\gamma}$

materials also exhibit a hysteresis loop which represents the A ferroelectric has been defined as a dielectric having a form. The structure of ferroelectric materials becomes less spontaneous polarization which can be reversed in sign. It therefore must have a polar structure with no center of symmetry. In changing the direction of the polar axis the structure must pass through an intermediate non-polar stage and distorted as the temperature increases and undistorted at and relation between magnetic induction $ar{B}$ and magnetic field H. the polar structure is a distortion of this more symmetrical

5.1. TERNARY PEROVSKITES

above a temperature called the Curie point.

Titanates

Barium titanate has the ideal cubic perovskite structure above the Curie point, but on cooling below this temperature the oxygen and titanium ions shift to form a tetragonal of barium titanate is cooled below 120°C it breaks up structure with the c-axis about 1% longer than the other two. At about 0°C the symmetry of the crystal becomes orthorhombic, and at -90°C trigonal (see Fig. 5.2). When a crystal into many domains with c-axes oriented perpendicularly.

thickness was postulated to be caused by a surface layer different from the bulk of the order of 10-4 cm on the Merz(9) found that the switching time required to change the domain configuration depended on the applied field and the size of the sample. This dependence of switching time on crystal. The switching time depends linearly on the thickness of the sample if the field is kept constant (see Fig. 5.3).

gonal strain in the surface did not vanish even when the rrystal was heated above the Curie temperature of the bulk. As ported by Känzig from electron diffraction experiments which indicated that there was a tetragonal strain in the further support for the postulation, Chynoweth reported that pyroelectric currents could be produced in single crystals of The presence of the surface layer on BaTiO, also was restrain in the bulk below the Curie temperature. (10) The tetrasurface layer which was slightly larger than the tetragonal

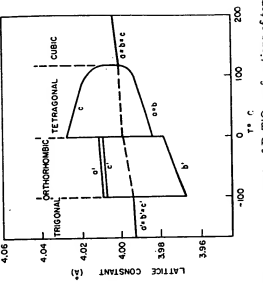
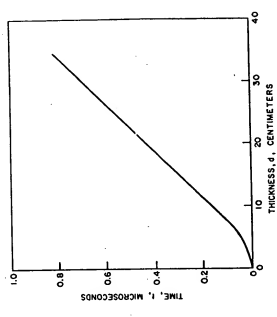


Fig. 5.2. Lattice constants of BaTiO, as functions of temperature (after H. F. Kay and P. Vousden, Phil. Mag. 40, 229 (1948)).



Fra. 5.3. Switching time for infinite field vs. thickness of the sample (after Merz⁽⁹⁾).

barium titanate above the Curie temperature even though no electric field was applied, (11) but in spite of these studies, there still appears to be considerable doubt as to the existence of the space-charge layer on the surface.

The tetragonal distortion in the structure of BaTiO₃ resulted in the formation of dipoles. Merz⁽¹²⁾ measured the dipole moment of BaTiO₃ single crystals and found it to be 18×10^{-6} coulombs/cm² at 120° C and 26×10^{-6} at ambient temperature. In addition, he⁽¹³⁾ also measured the dielectric constant and found that it was much greater perpendicular to the c-axis than along it (see Fig. 5.4).

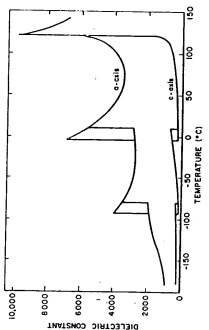


Fig. 5.4. Dielectric constant of BaTiO₃ as a function of temperature (after Merz⁽¹³⁾).

The dielectric constant of ceramic barium titanate was found to be about 1500, which falls between the values obtained along the c-axis and a-axis of the single crystal. The value decreases slowly with frequency to 10^6 c/s, and then decreases further to a value of 126 at 24×10^3 Mc. When the measurement is made under pressure, the dielectric constant increases⁽¹⁴⁾ and the Curie point shifts to lower temperatures.⁽¹⁵⁾

Because of this high dielectric constant, experimental studies have been conducted to improve the properties of barium titanate for energy converter and capacitor applications. A figure of merit for capacitor applications is de-

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noted as (time constant) RC which is the time for a capacitor to discharge to 1/e of the charging voltage. Time constants were measured for various grades of BaTiO₃ and for BaTiO₃ with different additives. The results are presented in Table

Table 5.1. Measurements on BaTiO₃ Compositions Fired I hr at 1450° C (after Hoh and Pirigy⁽¹⁶⁾)

RO in sec.	10 52 220 220 3,000 12,500 10,600 9,800 12,000 12,000 10,500 13,000
Additive	none none none 0.1 wt% Cr ₂ O ₃ 0.5 mole % UO ₂ as U ₃ O ₄ 0.5 mole % UO ₂ as U ₃ O ₄ 0.5 mole % UO ₂ as U ₃ O ₄ 0.5 mole % UO ₂ as U ₃ O ₄ 0.5 mole % UO ₂ as U ₃ O ₄ 0.5 mole % UO ₂ as U ₃ O ₄ 0.5 mole % UO ₂ as
BaTiO ₃ starting material grade	 C.P. Commercial capacitor Commercial piezoelectrio High purity Commercial capacitor High purity Gommercial capacitor Commercial capacitor

The results showed that the additives improved the time constant by several orders of magnitude. The compounds $\operatorname{Cr}_2\operatorname{O}_3$ and CaSnO_3 improved RC only at room temperature, but the solid fluoride additives showed the most promise.

MacChesney et al. (17) improved the stability of the properties with respect to temperature by adding La₂O₈. The temperature coefficient of capacitance and dissipation factor was reduced to low values by adding 1 mole % La₂O₃.

In other studies, increased stability of BaTiO3 has been

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which means that electric polarization takes place when it is Barium titanate also exhibits piezoelectric properties subjected to mechanical strain, and inversely the material This effect reverses in sign upon reversal of the electric field. This is in contrast to electrostriction exhibited by all dielecmechanically deforms upon application of an electric field. trics under an applied field.

 -310×10^{-8} stateoul/dyne for d_{31} where d_{33} is the proportionpendicular to its c-axis and d_{31} is the proportionality between Determination of the piezoelectric properties of singlecrystal BaTiO₃ gave 950×10^{-8} statcoul/dyne for d_{33} and ality between the charge developed on the two faces perthe charge on the same two faces and the force applied when the force is perpendicular to the c-axis. (21-24)

lites. The piezoelectric moduli measured for ceramics depend In order to obtain this effect in ceramic barium titanate, it must be first poled by d.c. voltages of $20-30~\mathrm{kV/cm}$ which line up approximately 10% of the c-axis of the crystalon the effectiveness of the poling operation.

c-axis and 0.47 Å for the Pb ions which were much larger displacements than those found in barium titanate (see Fig. 5.5). At 490°C, PbTiO₃ changes from a tetragonal form to a The results indicated a shift of 0.30 Å for the Ti ion along the tortion of the perovskite structure. Shirane and Pepinsky⁽²⁵⁾ studied the structure with X-ray and neutron diffraction. cubic form (see Fig. 5.6). The energy absorption at this trans-Lead titanate is also a ferroelectric with a tetragonal disformation temperature is 1150 cal/mol. (26)

Studies on single crystals indicate that the Curie point is at 495°C. The domain structures of these single crystals were reported to be essentially the same as that observed in The dielectric constant of PbTiO₃ is about 100 at room temperature and reaches a peak of about 1000 at 490°C.

cept for specimens containing 1 mole % CaF, which gave The piezoelectric coefficient d_{33} for PbTiO₃ with additives was found to be less than 30×10^{-12} coulombs/newton exvalues as high as 130×10^{-12} coulombs/newton. (23)

reported for materials with the following additives: 1% $NiZrO_3$, (Ti, Zr, Sn) $O_2 + Bi_2O_3$, 6% (Bi $_2O_3$, 2Zr O_2), (Zr, Sn) $O_2 + Li_2O_3$, (Zr, Sn) $O_2 + Al_2O_3$, 2.37(CaTi O_3) + 1.43(Bi $_2$ Sn O_5) + $+2.85 \mathrm{SnO}_2$, $1-25 (\mathrm{CaZrO_3})$, and $41 \mathrm{Fe_2O_3} + 10 \mathrm{TiO_2} + 9 \mathrm{FeO} +$ +5NiO + 5CoO.

bitanate after reduction and re-oxidation and with metal additives. However, it is probable that the high dielectric constants reported for these materials are high because of semi-High dielectric constants have been produced in barium conductivity of the samples or the presence of a thin nonconducting skin on a conducting medium.

curves were flatter with decreasing particle size. This is an important method of obtaining materials whose dielectric Khodakov (18) attempted to modify the properties of BaTiO₃ that the peaks in the dielectric constant versus temperature constants are high and relatively independent of temperaby using a fine particle size of BaTiO₃ $(1-20\mu)$ and found

constant as a function of temperature revealed that the sprayed barium titanate ceramic coatings. The coatings were predominantly the cubic phase. The metastable cubic phase transforms to the tetragonal phase by annealing the ceramic curves were much flatter than that for normal barium titanate. Blinton and Havell(19) studied the properties of flameat 1400°C for 2 hr in air or helium. A study of the dielectric

Interesting dielectric materials also were prepared by crystallizing BaTiO₃ with feldspar BaAl₂Si₂O₈ by heat-treating glasses having compositions corresponding to x BaTiO₃+ +(100-x) (BaAl₂Si₂O₈). Herczog⁽²⁰⁾ proposed the mechanism of crystallization as being

Glass
$$A \xrightarrow{600^{\circ} to} 10^{\circ}$$
 BaTiO_s(Cr)+glass B
Glass $B \xrightarrow{750^{\circ} to} BaAl_{\bullet}Si_{\bullet}O_{\bullet}(Cr) + BaTiO_{\circ}(Cr)$

stant was nearly independent of temperature, and materials ment were controlled the particle size could be varied from 0.01 to 1μ . The size of the particles were quite uniform for with a particle size of 1μ were found to have the highest If the heating rate and the final temperature of heat treatany treatment. For particles below 0.2μ the dielectric con-

Strontium titanate has the cubic perovskite structure, and is not ferroelectric at room temperature even though it has a dielectric constant of 200. There are conflicting reports on

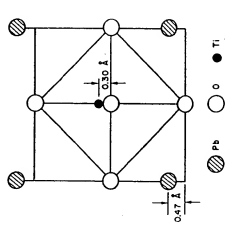


Fig. 5.5. Structure of PbTiO₃ (after G. Shirane, R. Pepinsky and G. B. Frazer, Acta Cryst. 9, 131 (1956))

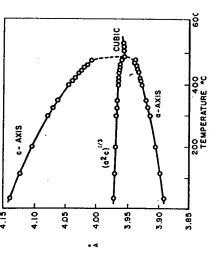


Fig. 5.6. Temperature dependence of volume and lattice constants for PbTiO₃ (after Shirane et al. (70)).

tures, even though Gränicher (20) did report the observation of the possibility of it being ferroelectric at very low temperaa hysteresis loop with a field of 300 V/cm at 4°K. The spon-

taneous polarization was reported to be 3×10^{-6} coulomb/ cm^2 and the remnant polarization 1×10^{-6} coulomb/cm².

cubic at 1000°C.(30) It has a room-temperature dielectric Calcium titanate has an orthorhombic structure at room temperature, the structure becomes tetragonal at 600°C and constant of 100, but it is not ferroelectric.

Niobates and Tantalates

of this group. The transition temperature on heating is Potassium niobate, KNbO₃, is the best-known compound -12°C for the rhombohedral to orthorhombic transforma-

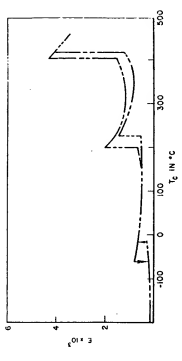


Fig. 5.7. Dielectric constant of KNbO₃ as a function of temperature (after Shirane et al.(311).

is about ten times as high as the value for BaTiO3. The values gent at these transitions is approximately 0.3.(31, 32) This tion, 412°C for the tetragonal to cubic transformation and -59°, 200° and 407°C on cooling (see Fig. 5.7). The loss tanof saturation polarization have been obtained from hysteresis loops and found to be 0.9×10^{-6} coulombs/cm² at room tion, 224°C for the orthorhombic to tetragonal transformatemperature.

tion of a strong field of the order of 10 kV/cm. Once ferroelectricity is induced, the crystals remain ferroelectric from Sodium niobate is not ferroelectric and may be antiferroelectric, but ferroelectricity can be induced by the applica--55°C to 200°C. Vousden⁽³³⁾ and Francombe⁽³⁴⁾ indicated FERROELECTRICITY

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that there are several transformations in NaNbO₃ at 20°, 390° , 420° , 560° and 640° C. The last is the tetragonal to cubic transformation. The temperature dependence of the dielectric constant is given in Fig. 5.8.

A rectangular hysteresis loop with the same saturation polarization value of BaTiO₃ was investigated by Matthias⁽³⁵⁾ who reported the Curie temperature for NaTaO₃ to be 475°C.

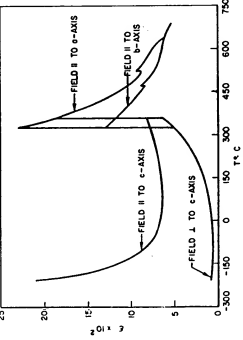


Fig. 5.8. Temperature dependence of dielectric constant of NaNbO₃ (after L. E. Cross and B. J. Nicholson⁽³¹⁾).

Potassium tantalate was reported as having a phase transformation between 10°K and 20°K and a hysteresis loop below 13°K⁽³⁶⁾. Smolenskii reported a Curie temperature of 247°C for RbTaO₃⁽³⁷⁾

Zirconates and Hafnates

Lead zirconate, PbZrO₃, is antiferroelectric, that is although there is a dipole moment in each unit cell, the arrangement of the moments in adjacent cells is such as to cause a net dipole moment of zero. At 230°C, the symmetry of the structure changes from orthorhombic to cubic (see Fig. 5.9). The temperature variation of the dielectric constant of lead zirconate is given in Fig. 5.10.

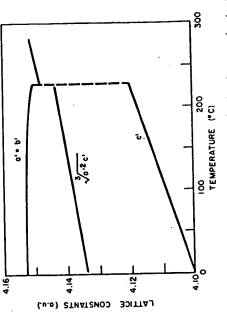


Fig. 5.9. Temperature variation of the lattice constants of lead zirconate (after E. Sawaguchi, J. Phys. Soc. Japan 8, 615 (1953)).

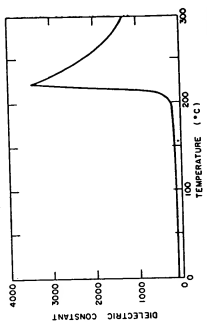


Fig. 5.10. Dielectric constant of lead zirconate as a function of temperature (after Roberts⁽⁸⁶⁾).

Lead hafnate transforms to a new structure at 160°C and transforms at 210°C to a cubic form. The two lower forms are antiferroelectric and the high-temperature form is paraelectric.(38)

5.2. Solid Solutions

BaTiO,-SrTiO,

One of the most widely studied solid solution systems is that between BaTiO, and SrTiO,. There is complete solid

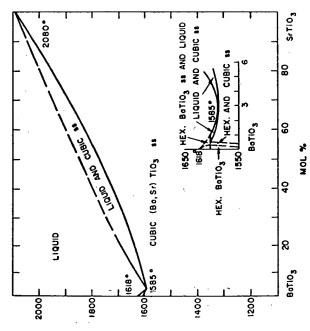


Fig. 5.11. Phase diagram for the BaTiO₃-SrTiO₃ system (after J. D. Basmajian and R. C. DeVries, J. Am. Geram. Soc. 40, 374 (1957)).

ing linearly with the substitution of Sr in $BaTiO_3^{(39,40)}$ (see Fig. 5.11). The Curie point also decreases with increasing solution in this system, with the size of the unit cells decreas-

30 mole % addition, and then decreases. It is interesting that the low-temperature Curie point materials do not show the nate at 10° cycles. (41-42) A minimum in the activation energy tric constant increases, reaches a maximum of 8000 at about decrease in dielectric constant that is found for barium titaamounts of strontium substitution (see Fig. 5.12). With the first additions of strontium the ambient temperature dielecobtained from d.c. measurements was found at

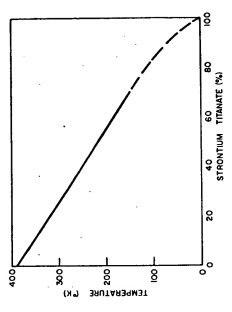
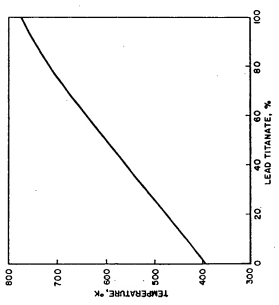


Fig. 5.12. Variation of Curie temperature as a function of composition, (Ba, Sr)TiO₂ (after Rushman et al. ⁽³⁹⁾).

BaTiO3-PbTiO3

ture (see Fig. 5.13),(34) decreases the d.c. resistivity and the Lead titanate also forms complete solid solution with barium titanate. The addition of lead titanate lowers the room temperature dielectric constant, increases the curie temperaactivation energy⁽⁴³⁾, and improves the piezoelectric properties of barium titanate. (44, 45)



Fro. 5.13. Variation of Curie temperature with composition, (Ba, Pb)TiO₂ (after G. Shirane, S. Hoshino and K. Suzuki, J. Phys. Soc. Japan 5, 456 (1950)).

BaTiO3-CaTiO3

Calcium titanate is soluble in BaTiO₃ up to 25 mole % and BaTiO₃ dissolves in CaTiO₃ to about the same extent (see Fig. 5.14). The addition of calcium to barium titanate lowers the room-temperature dielectric constant, and small amounts improve the piezoelectric properties.

BaTiO₃-(Ba, Sr, Ca)ZrO₃

The BaTiO₃-BaZrO₃ phase diagram is shown in Fig. 5.15. The addition of barium, strontium and calcium zirconate to barium titanate lowers the Curie point and broadens the maximum dielectric constant. (46)

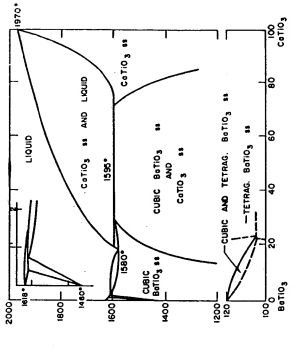


Fig. 5.14. Phase diagram for the BaTiO₃-CaTiO₃ system (after R. C. DeVries and R. Roy, J. Am. Geram. Soc. 38, 145 (1955)).

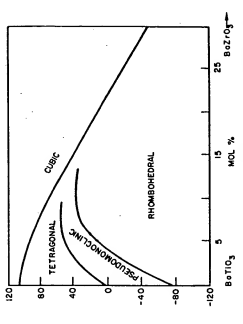


Fig. 5.15. Phase diagram for the system BaTiO₅-BaZrO₅ (after T. N. Verbitskaya, G. S. Zhdanov, Yu. N. Venevtsev and S. P. Solov'ev, *Kristallografitya*, 3, 189 (1958)).

$BaTiO_{a}$ - $BaSnO_{3}$

Dungan et al. '47) found that additions of BaSnO₃ to BaTiO₃ lowers the Curie temperature, and increases the unit cell size. Figure 5.16 shows the variation in Curie temperature with additions of BaSnO₃.

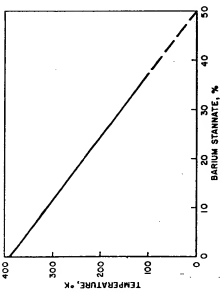


Fig. 5.16. Variation of Curie point of barium titanate with the addition of barium stannate (after Dungan et al. 47).

BaTiO,-BaHfO,

A study of the BaTiO₃-BaHfO₃ system by Fresenko and Prokopolo⁽⁴⁸⁾ showed that it was quite similar to those of BaTiO₃-BaZrO₃ and BaTiO₃-BaSnO₃. Payne and Tennery⁽⁴⁰⁾ made dielectric measurements and X-ray diffraction studies in this system and found that the dielectric constant for each sample increased as the BaHfO₃ concentration was increased to 16 mole % BaHfO₃ and then decreased with further BaHfO₃ additions. They suggest that the ferroelectric-paraclectric transition for the composition containing 16 mole % BaHfO₃ was of second order and occurred between a ferroelectric rhombohedral phase and a paraelectric cubic phase.

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Other BaTiO₃ Solid Solutions

More complex systems with barium titanate as one of the constituents have also been studied. The phase diagram for the system BaSnO₃-BaTiO₃PbSnO₃-PbTiO₃ is shown in Fig.

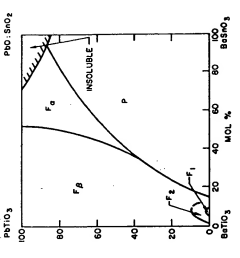
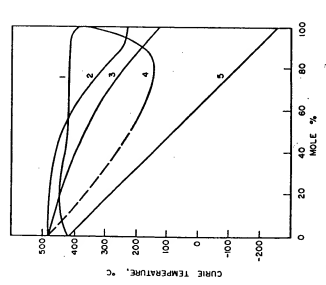


Fig. 5.17. Phase diagram for the PbTiO₃-BaTiO₃-PbO:SnO₃-BaSnO₃ system, P = paraelectric, cubic phase, F_a = ferroelectric, rhombohedral phase, F_g = ferroelectric, tetragonal phase, F_1 = ferroelectric, rhombohedral phase, F_2 = ferroelectric, orthorhombic phase (after T. Ikeda, J. Phys. Soc. Japan 14, 1292 (1869)).

PbTiO3-KNbO3

The phase diagram of the PbTiO₃-KNbO₃ system and dielectric properties of the solid solutions were determined by Tien *et al.*⁽⁸⁰⁾ Where solid solution was formed between compounds with one common cation, the Curie temperature varied more or less linearly with composition. Figure 5.18 presents this data for the KNbO₃-NaNbO₃, PbTiO₃-PbZrO₃, PbTiO₃-BaTiO₃, PbTiO₃-RNbO₃ systems. In the PbTiO₃-KNbO₃ system where no ions are

common the preparation of homogeneous specimens proved to be difficult. Tien et al. feel that the minimum that exists is common for all systems which involve ferroelectric or antiferroelectric compounds not containing a common ion, point-



(2) PhTiO,-PhZrO, (3) PhTiO,-BaTiO, (4) PbTiO,-NaNbO, (5) KNbO,-KTaO, (after Tien et al. (60) Fig. 5.18. Curie temperatures in perovskite systems: (1) KNbo,-NaNbo,

ing out that this is true of the BaTiO_s-PbZrO_3, NaNbO_s-NaNbO₃-PbTiO₃ as well as the PbTiO₃-KNbO₃ $Pb\bar{Z}rO_3$, system.

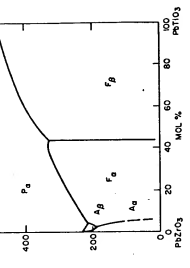
PbZrO₃-PbTiO₃

the antiferroelectric properties of PbZrO₃. Figure 5.19 shows a phase diagram of the PbZrO₃-PbTiO₃ system. The addition Addition of barium, strontium and titanium ions destroys of lead titanate to lead zirconate appears to lower the dielec-

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tric constant and increases the temperature of its maxi-

ing structure the induced piezoelectric effects are enhanced as the composition approaches the phase boundary. In the system lead titanate-lead zirconate, lead titanate-lead oxide: tin oxide, lead zirconate-lead oxide: tin oxide and the Jaffe et al. (52) found that in systems where compositional boundaries exist between ferroelectric phases of slightly differead titanate-lead hafnate, one of the compositions 45%



 $F_{\beta} = \text{ferroelectric}$, tetragonal phase P = paraelectric, cubic phase, $A_{\alpha} = \text{antiferroelectric}$, ortho-Fig. 5.19. Phase diagram for the PbTiO,-PbZrO, system, $A_{\beta} = \text{antiferroelectric}, F_{\alpha} = \text{ferroelectric},$ (after E. Sawaguchi, J. Phys. Soc. Japan 8, 616 (1953)). rhombohedral phase, rhombic phase,

a radial coupling coefficient of 0.3 at 275°C which is twice PbTiO₃-55% PbZrO₃ has a Curie temperature of 340°C and that of barium titanate. Another containing 47.25% PbTiO₃, of all compositions studied—74 22.75% PbZrO₃ and 30% PbO:SnO₂ had the highest piezo-×10-12 coulomb/newton. electric coefficient (d₃₁)

electric performance. In more detailed studies Ikeda^(s4) showed that improved piezoelectric ceramics were obtained when Pb(Ti,Zr)O₃ was modified by the addition of A¹⁺B⁵⁺O₃ $\mathrm{Zr})\mathrm{O}_3$ ceramics near the phase boundary improved the piezo-Ikeda (53) reported that the addition of LaFeO to Pb(Ti $(A = K, Na, B = Sb, Bi) \text{ or } A^{3+}B^{3+}O_3 (A = Bi, La; B = Fe, Al$

Cr). Dielectric constants above 1500 and radial coupling coefficients above 0.6 were obtained with Na+ and Sb5+ substitution.

PbZrO₃-BaZrO₃

mum of lead zirconate is shifted to lower temperatures. (55) When barium replaces lead in PbZrO3, the dielectric maxi-In addition, the maximum dielectric constant becomes great-

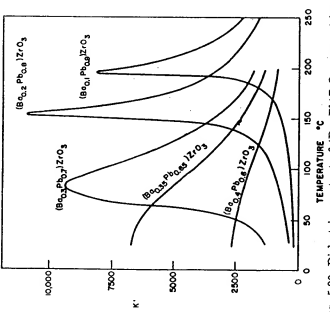


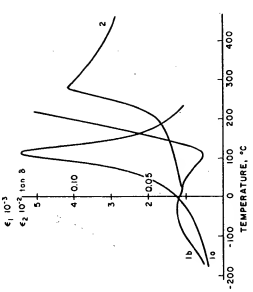
Fig. 5.20. Dielectric constant of (Ba, Pb)ZrO, compositions at varying temperatures (after Roberts⁽⁶⁵⁾)

er and the high values are maintained over a broader range mum is lowered, but the temperature range of high dielectric constant values become broader. The dielectric constant of Bao.35Pbo.85ZrO3 stays above 6000 from room temperature of temperature (see Fig. 5.20). The largest value is obtained for a composition of (Ba_{0.20}Pb_{0.80})ZrO₃ and when still larger percentages of barium are substituted, the dielectric maxiup to 60°C making it useful for capacitor application.

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5.3. COMPLEX PEROVSKITES

ported anomalies in dielectric constants at temperatures of Viskov et al. (56) reported that the compounds Ba(Bio.s Nbo.s)O3, Ba(Bio.sTao.s)O3, Ba(Bio.sUo.s)O3, Ba(Bio.s7Wo.ss)O3 and Ba(Bio.s Moo.s)O3 had distorted unit cells and high dielectric constants which peaked with temperature. They re-420°, 410°, 320°, 400° and 260°C for the compounds con-



and $Pb(X_0, _\delta N\tilde{b}_0, _\delta)O_3$; (i.e. and 1b) ε_1 and $\tan \delta$ for $Pb(Fe_0, _\delta Nb_0, _\delta)O_3$; (2) ε_2 for $Pb(Y_0, _\delta Nb_0, _\delta)O_8$ (after Smolenskii et al., ε_1). Fig. 5.21. Temperature dependence of ϵ for Pb(Fe_{0.5}NP_{0.5})O₈

taining Nb, Ta, V, W and Mo, respectively, and felt that the first three were ferroelectric and the last two antiferroelectric. However, evidence for this assumption was lacking.

pacts that Pb(Fe_{0.5}Nb_{0.5})O₃ and Pb(Yb_{0.5}Nb_{0.5})O₃ might be Smolenskii et al. (57) found from studies on powder comerroelectrics with Curie temperatures of 112°C and 280°C, respectively. The temperature dependence of permittivity and loss tangent for these compounds are shown in Fig. 5.21. The compound Pb(Feo.sNbo.s)O3 exhibited a hysteresis loop, out Pb(Yb_{0.5}Nb_{0.5})O₃ only showed a maximum in its permitivity and therefore may be antiferroelectric.

permittivity peak was observed at 38°C in the dielectric constant—temperature curve for Pb/Mr, W., 10... The com-

permittivity peak was observed at 30 C in the dielectric constant—temperature curve for $Pb(Mg_{0.5}W_{0.5})O_3$. The compounds $Pb(Fe_{0.67}W_{0.33})O_3$ and $Pb(Fe_{0.5}Ta_{0.5})O_3$ exhibited hysteresis loops at liquid-oxygen temperature and the latter compound exhibited a maximum in its dielectric constant at -30° C.

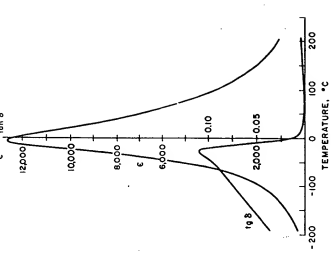


Fig. 5.23. Temperature dependence of ε and $\tan \delta$ values for Pb(Mg_{0.88}Nb_{0.67})O₈ (1 kc) (after Smolenskii εt $al.^{(62)}$).

Single crystals of $Pb(Ni_{0.33}Nb_{0.67})O_3$ were prepared by Myl'nikova and Bokov and reported to be ferroelectric. A maximum was observed in a plot of its dielectric constant versus temperature, and a hysteresis loop was observed at a temperature of $-196~{}^{\circ}C.^{(62)}$

Smolenskii and Agranovskaya⁽⁶³⁾ studied a large number of perovskite-type compounds and found two new ferroelectric materials, Pb(Mg_{0.33}Nb_{0.67})O₃ and Pb(Ni_{0.33}Nb_{0.67})O₃. Figure 5.23 shows the temperature dependence of the dielec-

Smolenskii et al.(39) also found that Pb(Sc_{0.5}Nb_{0.5})O₃ and Pb(Sc_{0.5}Ta_{0.5})O₃ were ferroelectric materials. The dielectric constant of the niobium- and tantalum-containing compounds exhibited a maximum at 90° and 20°C, respectively (see Fig. 5.22). The characteristic drop in the loss tangent corresponds to a maximum of the dielectric constant. Hysteresis loops were obtained for both compounds at 18°C. The spontaneous polarization for Pb(Sc_{0.5}Nb_{0.5})O₃ at 18° equaled 3.6 microcoulombs and the coercive force 6 kV/cm.

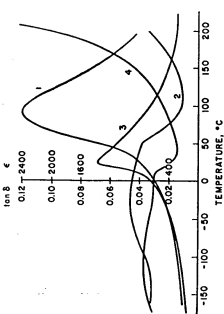


Fig. 5.22. Temperature dependence of ε and $\tan \delta$ of Pb(Sc_{0.5}Nb_{0.5})O₃ and Pb(Sc_{0.5}Ta_{0.5})O₃: (1, 2) ε and $\tan \delta$ of Pb(Sc_{0.5}Nb_{0.6})O₃; (3, 4) ε and $\tan \delta$ of Pb(Sc_{0.5}Nb_{0.6})O₃; (after Smolenskii ε t $al.^{(68)}$).

Maxima in dielectric constant with temperatures were observed for Pb(Lu_{0.5}Nb_{0.5})O₃, Pb(Yb_{0.5}Ta_{0.5})O₃ and Pb(In_{0.5}Nb_{0.5})O₃ at 280° for the first two compounds and 90° for the last, but they were probably antiferroelectric. (50) Studies on single crystals of Pb(Co_{0.5}W_{0.5})O₃ (60) showed that it had a maximum in its dielectric constant at 32°C and exhibited a double hysteresis-loop characteristic of an antiferroelectric material.

Measurements on $Pb(Fe_{0.67}W_{0.38})O_3$ and $Pb(Fe_{0.5}Ta_{0.5})O_3$ indicate that these compounds are true ferroelectrics, while $Pb(Mg_{0.5}W_{0.5})O_3$ appears to be antiferroelectric.⁽⁶¹⁾ A sharp

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tric constant and loss tangent for Pb(Mg_{0.33}Nb_{0.67})O₃. The dielectric constant reached a maximum value of 12,600 at -15° C when a frequency of 1 kc/s was used in the measurements. A hysteresis loop was observed at -130° C and a spontaneous polarization value of 14×10^{-6} coulombs was calculated. The compound Pb(Ni_{0.33}Nb_{0.67})O₃ also appears to be a ferroelectric material.

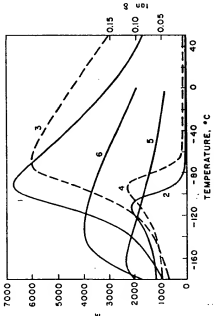


Fig. 5.24. The temperature dependences of ε and $\tan \delta$: (1, 2) ε and $\tan \delta$ for single crystals $Pb(Mg_{0.38}Ta_{0.67})O_8$ (1 kc); (3, 4) ε and $\tan \delta$ for single crystals $Pb(Vo_{0.38}Nb_{0.67})O_8$ (1 kc); (5) ε for single-crystal $Pb(Ni_{0.38}Ta_{0.67})O_8$ (450 kc), (6) ε for single-crystal $Pb(Co_{0.38}Ta_{0.67})O_8$ (1 kc), (after Bokov εt $aL^{(64)}$).

Bokov and Myl'nikova'⁽⁴⁴⁾ prepared single crystals of compounds Pb(Ni_{0.33}Ta_{0.67})O₃, Pb(Mg_{0.33}Ta_{0.67})O₃, Pb(Co_{0.33}Nb_{0.67})O₃, Paxx = 28 kV/cm, at $t = -150^{\circ}$ C, for Pb(Co_{0.33}Nb_{0.67})O₃,

 $E_{\text{max}} = 38 \text{ kV}/\text{cm}$, at $t = 20^{\circ}\text{C}$, for Pb(Mg_{0.33}Ta_{0.67})O₃, $E_{\text{max}} = 45 \text{ kV}/\text{cm}$, at $t = 182^{\circ}\text{C}$, for Pb(Co_{0.33}Ta_{0.67})O₃, $E_{\text{max}} = 70 \text{ kV}/\text{cm}$, at $t = -196^{\circ}\text{C}$, and for Pb(Ni_{0.33}Ta_{0.67})O₃, $E_{\text{max}} = 150 \text{ kV}/\text{cm}$, at $t = -196^{\circ}\text{C}$. The Curie temperatures of compounds Pb(Mg_{0.33}Nb_{0.67})O₃, Pb(Mg_{0.33}Ta_{0.67})O₃, Pb(Co_{0.33}Nb_{0.67})O₃, Pb(Ni_{0.33}Nb_{0.67})O₃, Pb(Ni_{0.33}Ta_{0.67})O₃, Pb(Ni_{0.33}Ta_{0.67})O₃, Pb(Ni_{0.33}Ta_{0.67})O₃, Pb(Ni_{0.33}Ta_{0.67})O₃, Pb(Ni_{0.33}Ta_{0.67})O₃, Pb(Ni_{0.33}Nb_{0.67})O₃, Pb(Ni₀

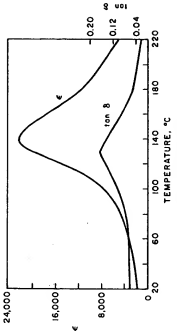


Fig. 5.25. The temperature dependence of ε and $\tan \delta$ of single crystal Pb(Zn_{0.33}Nb_{0.87})O₃ (1 kc) (after Bokov *et al.*⁽⁶⁴)).

Johnson et al. (65) conducted an extensive study of complex ferroelectric solid solutions $Pb(Sc_{0.5}Nb_{0.5})_x \phi_{1-x}O_3$ where ϕ is either Ti, Zr, or Hf, finding that the maximum value of the spontaneous polarization of a system decreased as Ti was replaced by Zr and Zr by Hf and the Curie temperature increased as ϕ increased. In addition, the Curie points decreased in value as Ti was replaced by Zr and Zr by Hf. A complete list of ferroelectric compounds, Curie points and polarization values are given in Table 5.2.

5.4. EFFECT OF NUCLEAR IRRADIATION

In recent years there has been considerable interest in the effect of nuclear radiation on the properties of ferroelectric materials. In one study, Glower and Hester⁽⁶⁰⁾ found that

Table 5.2. Ferroelectrics Data

-	T °C	Ps (at T °C)	Refs.
BaTiO ₃ .	120	10-6	69
		26.0 (23)	3
Portio	490	750.0 (23)	20
KNDO3	435	0.9(23)	35
		30 (250)	
KTaO3	-260	•	35
Na.NbO3	-200		7.1
CdTiO	-218		72
Pb(Cd _{0.8} W _{0.8})O ₃	-		73
Pb(Sco. 8Nbo. 8)O3	90	3.6 (18)	5.0
Pb(Sc., Tao., s)O3	26	•	58
Pb(Fe _{0.67} W _{0.33})O ₃	-75		74
Pb(Fe.sNb.s)O3	112		75
Pb(Feo. 8 Tao. 8)O3	-30		92
Pb(Mg _{0.33} Nb _{0.67})O ₃	-12	14 (-130)	64
Fb(N10.33Nb0.67)O3	-120		64
FO(N10.33 T80.67) O3	-180		64
Fb(Mg _{0.33} Ta _{0.67})O ₃	86-		64
F D (Co. 33 N Do. 67) O3	-70		64
F0(C0, 33 1 80, 67) U3 Ph(Zn - Nh) O	-140		64
- C(2110.3311 C0.87) C3	140		64

nuclear reactor irradiation of single crystal of BaTiO₃ produced increases in the coercive field (E_c) and a decrease in the remnant polarization (P_r) . Crystals in the polarized state during irradiation were more resistant to radiation damage than were virgin crystals and the radiation damage rate was only slightly dependent upon crystal thickness. In their conclusions, they interpreted the changes in E_c and P_r in terms of a radiation model involving a build-up of a space charge due to the trapping of ionized carriers in the domain walls of the crystals.

Hilczer⁽⁶⁷⁾ reported that irradiation of barium titanate with 10¹⁹ neutrons/cm² reduced the dielectric constant by up to 40%. Doses of 10¹⁴–10¹⁹ neutrons/cm² of (a) pile neutrons containing 10% fast neutrons; and (b) fast neutrons only that had passed through a 0.4-mm Cd foil gave the same effect, implying that fast neutrons produced the damage.

Schenk⁶⁸⁾ found that tetragonal BaTiO₃ when irradiated with neutron doses of 4.2×10^{18} neutrons/cm² at 35°C transformed to cubic with radiation damage. The lattice expanded 2.26% for a and 1.17% for c.

5.5. APPLICATIONS OF FERROELECTRIC MATERIALS

Use of Ferroelectric Properties

The high dielectric constants and ferroelectric behavior of perovskite-type compounds are probably the most important properties they exhibit.

Materials such as barium titanate cannot be used as capacitors in tuned circuits or filters where high-frequency stability is needed. However, they can be used as by-pass, blocking and smoothing capacitors which present a low-impedance path to an alternating current above a certain frequency.

The room-temperature dielectric constant of barium titanate can be raised by the addition of strontium titanate which lowers the Curie temperature. However, a flattening of the permittivity peak by using mixtures of alkali titanates and zirconates is more important. To date most of these ceramic capacitors have been used at low voltages as by-pass capacitors.

In addition, the hysteresis loop of ferroelectric single crystals makes them of potential use for information storage in electronic computors. Ferroelectrics also have been used as dielectric amplifiers. These are analogous to magnetic amplifiers, which require magnetic materials with narrow rectangular hysteresis loops.

The International Telephone and Telegraph Corporation recently reported a new use for ferroelectrics. This method for producing high-voltage a.c. or d.c. power was based on the fact that the dielectric constant of a ferroelectric is sensitive to temperature at the Curie point. The capacitor is held at the Curie point and then heated, lowering the dielectric constant. Since the charge cannot decrease because of a diode in the circuit, there must be a rise in capacitor voltage. This increased voltage also means an increase of electrical energy, thus there is a conversion of heat energy into elec-

trical energy. This scheme has been proposed for use in a space vehicle which spins so it alternately faces toward and away from the sun.

Use of Piezoelectric Properties

One of the best known uses of piezoelectric materials is for the measurement of force or pressure. However, since there is a current leakage with time, they have best been used for measuring dynamic pressures, in blast gauges and acceerometers. Recently, the Spark Pump, the heart of which was introduced by Clevite as an ignition source for gasoline engines. These elements were capable of achieving voltages is two lead zirconate-lead titanate piezoelectric elements, of 20,000 volts when mechanical pressure was applied.

Another use of piezoelectrics is the phonograph pick-up graph needle to an electrical signal. One of the methods of accomplishing this involves the use of two piezoelectric plates combined into a sandwich which is subjected to bending which transforms the mechanical energy from the phono-

Piezoelectric transducers have also been used for sound transmission and reception, and ultrasonic cleaning devices.

Quartz has been used for electric frequency control. The piezoelectric coupling causes a reaction with an electric drivng circuit which forces the circuit to oscillate at an exact frequency.

In a wave filter application, the impedance property of a orystal near a resonance point is used to allow passage of an electric signal which falls within a prescribed band of frequencies, while other frequencies are not passed.

The selection of materials for these applications depends on their piezoelectric constant. Some constants are listed in energy stored and the mechanical energy applied. A high coupling coefficient, that is the ability to convert from one Fig. 5.26 and the units in Table 5.3 for perovskite-type ceramics. Note that the k, electromechanical coupling coefficient, values show the relationship between the mechanical energy form of energy to another, is desirable in most of the applistored and the electrical energy applied or the electrical cations.

FERROBLECTRICITY

PIEZOELECTRIC CERAMICS-BASIC ACTIONS

	ELECTRUMECHANICAL EFFECTS	AND DIELECTRIC CONSTANTS
PLATES, BARS	THICKNESS EXPANDER POLED FIELD STRAIN	^d 33, ⁹ 33, ^k 33 ^Y 33, p
	LENGTH EXPANDER	M 40
	POLED FIELD STRAIN	in 15 4 11 1
`2	-	K3
DISCS	RADIAL EXPANDER	
m.	POLED FIELD STRAIN	da, 931, hp
		a E
	Š	d33, 933, k33
),	POLEO FIELO	Y33, p
		K3
SHEAR PLATE	SHEAR STRAIN	415, 915, ⁸ 15
Ţ		Y44. Y55, p
		ř.
TUBES	THICKNESS EXPANDER	
	×.	d33, 933, k33
		93,6
	LENGTH (Circumference EXPANDER Expander)	d31, 931, k31
		Y _{(1, p}
•		n

Fig. 5.26. Piezoelectric ceramics—basic actions (after Bulletin 9247 (1962), Clevite Corp.).

duction. Other constants are K, relative dielectric constant The constant d is the ratio of the strain developed to the age production and a high g constant for high voltage pron the material to space, and N, the frequency constant applied field or the short circuit charge density to the applied stress. The constant g is the ratio of the open circuit field charge density. A high d constant is desired for high amperto the applied stress or the strain developed to the applied which is the controlling dimension times the resonant frequency.

PREPARATION OF PEROVSKITE-TYPE COMPOUNDS

Table 5.3. Ceramic Properties Definitions (after Bulletin 9247 (1962), Clevite Corp.)

Property	Definition	MKS units
Electro- mechanical coupling coefficient k	mechanical energy stored electrical energy applied electrical energy stored mechanical energy applied	
Piezoelectric constants d	strain developed applied field short circuit charge density applied stress	meter/meter volts/meter coulombs/meter²
. 8	open circuit field applied stress strain developed applied charge density	volts/meter newtons/meters meter/meter coulombs/meter²
Relative dielectric constant K	e(permittivity of material) (permittivity of space)	
Modulus of elasticity. Y	stress strain	newtons/meter ² meter/meter
Density $\hat{\varrho}$		kg meters³
Frequency constant N	Controlling dimension Xresonant frequency	cps-meters

respectively, and the subscripts 4, 5 and 6 represent a double subscript which stands for a plane. For example, 4 represents the yz-plane, 5 represents the xz-plane and 6 represents the xy-plane. The first subscript describes the direction of the The subscripts 1, 2 and 3 indicate the x-, y- and z-axes, field, the second the direction of the strain. For K the sub-

script p means planar coupling. In the constant Y, the first subscript refers to the direction of stress and the second to the direction of the strain. The erties of several ceramic compositions. The PZT materials are special commercial compositions of lead zirconate-lead titanate solid solutions. Using this data, the best materials subscript for the dielectric constant refers only to the field. Table 5.4 presents the elastic, piezoelectric and dielectric propfor a particular application can be selected.

5.6. Theories of Ferroelectricity

the excellent treatment of this subject by Jona and Shirane Because of the importance of perovskite-type compounds erty is presented. For more details, the original papers or as ferroelectrics, a brief review of theories which have been proposed to explain the phenomena associated with this propin a book entitled Ferroelectric Crystals, published by Macmillan Company of New York, can be consulted.

BaTiO3, 776, 77) He assumes that BaTiO3 in all forms can be equation can be written as a series involving powers of the free condition and have subscripts X. Devonshire gives the which can be expressed as a function of temperature, stress and polarization. If the stress is initially taken as zero, an polarization P. The coefficients are functions of the stressequation with separate terms and coefficients for the compo-Devonshire proposed a phenomenological theory for considered to be a strained cubic crystal with a free energy nents of polarization in the axial directions x, y, z. The equa-

$$G_1 - G_{10} = \frac{1}{2} A^X (P_x^2 + P_y^2 + P_z^2) + \frac{1}{4} B^X (P_x^4 + P_y^4 + P_z^4) + \frac{1}{6} C^X (P_x^6 + P_y^6 + P_z^6) + \frac{1}{2} D (P_y^2 P_z^2 + P_z^2 P_z^2 + P_z^2 P_y^2),$$

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Table 5.4. Blastic, Piezoelectric and Dielectric Properties of Several Ceramic Compositions (after D. Berlincourt, B. Jaffe, H. Jaffe, and H. H. A. Krueger, IRE Nat'l Convention (1959),

	96w % BaTiO ₃ 6w % CaTiO ₃	PZT-4	PZT-6
Coupling coefficients			
k_{33}	0.49		0.675
, , , , , , , , , , , , , , , , , , ,	0.325		0.54
7.31 K:*	0.19	0.31	0.32
Piezoelectric	9 F.		9000
constants			
d_{33}	150	256	320
d_{31}	-68	-111	-140
d_{18}	257	450	495
Free dielectric			
constants			
K_1	1280	1360	1285
, K	1200	1200	1600
Frequency			
constants			
N ₁	2290	1650	1600
	2840	2000	1890
Lussuc			
e u	-	1	
ii	11.6	8.15	6.75
1/82 = YE	11.1	6.7	5.85
H	4.4	5.6	2.0
Density	5.5	7.6	7.6
Mechanical Q	200	900	75
Curie point	116 °C	340°C	3e0 °C
		_	

where G_{10} is the value of G_1 for the unpolarized, unstressed between the components of polarization along the x-, y- and crystal and the last term indicates that there is an interaction z-axis. From the equation given above, four sets of solutions which may correspond to minima in free energy can be obtained:

$$P_x = P_y = P_z = 0;$$

 $P_x = P_y = 0, P_z \neq 0$
 $P_z = 0, P_x = P_y \neq 0$
 $P_x = P_y = P_z \neq 0$

These represent the cubic phase, in which the polarization is clinic axes) and the rhombohedral forms respectively. The relative depths of the minima of the free energy function change with the coefficient A'. If this decreases steadily and zero, and the tetragonal, orthorhombic (referred to monoconstant values are chosen for the coefficients, the temperatures at which the transitions occur are those actually observed.

On determining the constants, Devonshire drew theoreti-Qualitative agreement between calculated and experimental cal curves for the spontaneous polarization, the free energy and the dielectric constants, over a range of temperatures. data was quite good.

of clamping on the nature of the transition. Probably one neous strain for a clamped crystal and discussed the effect of the most important conclusions that can be made from crystal would be of the second order, even though that of the use of these equations is that the transition of the clamped Devonshire also gave equations for calculating spontathe free crystal is of the first order.

culated. For example, the entropy change at the transition Using Devonshire's approach of determining the coefficients for his equations from certain properties and employing them to predict others, a number of quantities were calwas determined for $BaTiO_3$ and $\check{K}NbO_3$, in reasonable agreement with experimental data.

Using a model approach, Mason and Matthias⁽⁷⁸⁾ suggest is not at the center of the oxygen octahedra. Instead it is at ments from the center toward the exygen ions. When the have a dipole moment. However, if any dipole theory were that the stable position for the Ti4+ ion in barium titanate any of the six positions which correspond to slight displace-Ii4+ ion was in any of these positions the unit cell would correct, a number of polar liquids would be ferroelectrics which is not the case. In addition, with this theory it is not

possible to obtain good agreement with experimental calcu-

does not require the attribution of dipole moments to atomic displacement. (80) Only the electronic states of the TiO, octahedra are considered. The theory was adequate for deterplaced rather than titanium ions(79) and also a theory which Jaynes proposed a model in which oxygen ions are dismining the entropy change, but it predicts an infrared absorption line at $\sim 10\mu$, which was not detected.

Devonshire's model theory considers the dipole of an atom vibrating in the field of the neighbors. The dipole moment is not fixed in magnitude, but depends on the displacement from the equilibrium position.

However, in addition, he assumes that each atom has an electronic polarization and titanium also has an ionic polarization. It predicts that the direction of spontaneous polarization is along the z-axis, but this is a disadvantage when Slater's theory(81) is similar to Devonshire's model theory. it is applied to some structures.

emphasis on the covalent bond character in the occurrence of ferroelectricity. The problem with Megaw's theory is that the origin of ferroelectricity is sought in abrupt changes in the character of the bonds at each transition. Like all of the theoelectricity are associated with the off-center position of a highvalency cation in an octahedron. Megaw^{682, 83)} addèd the ries described above, the structural approach has its limi-In a structural approach, ferroelectricity and antiferro-

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9 CHAPTER

PHASE TRANSITION

ten associated with a change in ferroelectric properties. Some ter, an attempt will be made to describe phase transitions reported for perovskite-type compounds, whether or not they involve a ferroelectric transition. These phase transitions can be divided into those of the first kind which are associated with the absorption or liberation of heat along with discontinuous changes in entropy and in lattice parameters, and those of the second kind which involve a peak in heat capacity, in the coefficient of thermal expansion and in compressibility. A list of perovskite-type compounds with transition THE phase transitions in perovskite-type compounds are ofof these were described in the previous chapter. In this chaptemperatures and references is given in Table 6.1.

6.1. TERNARY PEROVSKITES

and below this temperature the tetragonal ferroelectric phase phase is formed, which has a unit cell with orthorhombic symmetry and still is ferroelectric with the direction of sponbic (110) directions. At -90°C another transition occurs of these phases is shown in Fig. 5.2. The volume changes at The phase transitions in barium titanate are probably the appears, and remains stable to about 5°C. Below 5°C a new taneous polarization being parallel to one of the original cu-The polar axis lies along one of the original cubic $\langle 111 \rangle$ directions. The thermal expansion of the cell parameters in each the phase transitions with rising temperature are 0.0006, best characterized. The cubic phase is stable down to 120°C, and the symmetry of the structure becomes rhombohedral.

Table 6.1. Phase Transitions

Compound	Transition temp., °C (to cubic at highest temp. listed)	Refer- ences
AgNbO, AgTaO, BaTiO, KNbO,	325, 550 485 -90, 5, 120 -10, 225,	A C
KTaO3 NaNbO3		
NaTaO ₃ PbHfO ₃ PbTiO ₃ PbZrO ₃	230	5, 6 4, 7 8 9 10, 11,
$SrTiO_3$	- 220	7, 12,
Pb(Co _{0.6} W _{0.6})O ₃ Pb(Cd _{0.6} W _{0.6})O ₃ Pb(Lu _{0.5} Nb _{0.8})O ₃ Pb(Lu _{0.5} Ta _{0.5})O ₃ Pb(Yb _{0.6} Nb _{0.8})O ₃	32 410 280 280 280	13, 16, 16, 17, 14, 14,
Sr(Co _{0.5} Mo _{0.5})O ₃ Sr(Co _{0.5} Wo _{.6})O ₃ Sr(Fe _{0.5} Nb _{0.5})O ₃ Sr(Fe _{0.5} Ta _{0.5})O ₃	320 400 250 250	20 20 20 20 20 20 20
$\begin{array}{c} { m Sr(Ni_{0.5}Mo_{0.5})O_{3}} \\ { m Sr(Ni_{0.5}W_{0.5})O_{3}} \\ { m Sr(Y_{0.5}Nb_{0.5})O_{3}} \end{array}$	300 830 630	19 19 21
$Sr(Y_{0.5}Ta_{0.5})O_{3}$ $Sr(Zn_{0.6}Mo_{0.5})O_{3}$ $Sr(Zn_{0.6}W_{0.5})O_{3}$	640 420	21 19
O1 (2110.5 W 0.5) U3	7 0 0	A.T

0.014 and 0.062 Å³ and the transition heats are 8, 22 and 50 cal/mole.

The cubic to tetragonal transition in barium titanate also is characterized by the appearance of domain patterns. Observation of ferroelectric domains between crossed nicols is an excellent method of studying phase transitions in ferro-

PHASE TRANSITIONS

electrostatically charged particles have been deposited on the surface are other methods which have been used to inelectric materials. Examining crystals after etching and after vestigate the domain pattern. Studies on tetragonal barium titanate using a polarizing microscope have shown that it is possible for domains to be polarized at 90° to each other. When the polar axis is perpendicular to the plane of a (001) plate, the domain is called a the domain is called an "a" domain. These domains are easily "c" domain, and when it lies within the plane of the plate, observed under polarized light, and their appearance or disappearance indicates the cubic to tetragonal or tetragonal to cubic transition.

If domains are polarized antiparallel to each other, they are called 180° domains and a field must be applied perpendicular to the polar axis to make them visible.

with a heat of transition of 440 cal/mole and PbHfO₃ shows type compounds, the structure of strontium titanate becomes cubic at -220°C and that of calcium titanate changes to cubic at 1260°C. Lead titanate has a transition from a tetragonal ferroelectric phase to a cubic phase at 490°C, Fig. 5.6, the pseudotetragonal antiferroelectric PbZrO₃ phase undergoes a first-order phase transition to cubic at 230°C (Fig. 5.9) transitions at 163°C and 215°C. The transition for PbHfO₃ at 215°C is probably an antiferroelectric transformation to Considering phase transitions involving other A²⁺B⁴⁺O₃cubic symmetry.

see Fig. 5.7, and the c/a for the unit cell of tetragonal KNbO₃ The phase transitions at 435°C, 225°C and -10°C in that the spontaneous strain in KNbO₃ is larger in all three phases. The transition energies are 190, 85 and 32 cal/mole, is 1.017 compared with the value of 1.010 for that of BaTiO₃. All of the phase changes are first order and exhibit temperature hysteresis. The structure of KTaO3 changes to cubic KNbO3 are quite similar to those found for BaTiO3 except symmetry at -260°C.

NaNbO₃ has orthorhombic symmetry. This antiferroelectric is ferroelectric. Above this temperature the structure of phase changes to a nonpolar pseudotetragonal phase at The structure of NaNbO₃ is monoclinic below -200°C and

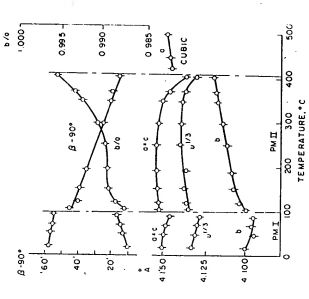
354°C. At 562°C the structure of NaNbO₃ becomes tetragonal, and cubic at 640°C. At room temperature, a polar structure can be induced by the application of a field and a double

A transition at 480° to cubic symmetry was found for NaTaO3; however, no anomaly in the dielectric constant has hysteresis loop can be observed. been detected.

at 325°C and from tetragonal to cubic symmetry at 550°C. forms from orthorhombic symmetry to tetragonal symmetry X-ray studies on AgNbO3 indicate that the structure trans-AgTaO3 shows similar phase transitions at 370°C and 485°C.

6.2. COMPLEX PEROVSKITE-TYPE COMPOUNDS

phase transitions. However, very few such transitions have type compounds listed in Table 5.2 must be associated with The onset of ferroelectricity in the complex perovskite-



Pb(Cdo., Wo.,)O, as functions of temperature (after Rogin-Fig. 6.1. Parameters of the perovskite pseudocell of skaya et al.(16)].

been reported with the ferroelectric data. In some cases, the to prove that small distortions existed in the ferroelectric back reflections in the X-ray patterns have been too poor phases.

have been studied. A pseudomonoclinic form of ${
m Pb}({
m Cd}_{0.5}{
m W}_{0.5}){
m O}_3$ has been found to exist up to 100°C, at which temperature Phase transitions for a few complex perovskite compounds it transforms to a second monoclinic form and at 410°C it becomes cubic (see Fig. 6.1). The cubic form has the ordered

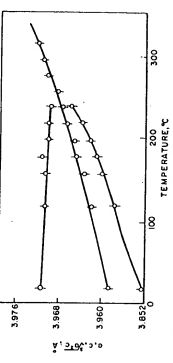


Fig. 6.2. Parameters of $\mathrm{Sr}(\mathrm{Fe_{0.6}Nb_{0.6}})\mathrm{O_3}$ as a function of temperature (after Kupriyanov et al.(17)).

equal to 8 Å. The authors feel that the monoclinic form of perovskite structure with a unit cell of "a" approximately Pb(Cd_{0.5}W_{0.5})O₃ below 100°C is antiferroelectric.

The phase transformation of Sr(Fe_{0.5}Nb_{0.5})O₃ involves a 6.2). From the unit cell size given there does not appear to be any ordering of the B ions in this compound. A similar phase change from a tetragonal to cubic form at 220°C (see Fig. transition was reported for Sr(Fe_{0.5}Ta_{0.5})O₃.

The compound Sr(Yo.s Tao.s)O3 was found to transform from a rhombohedral form to a cubic form at 640°C, similar to that in $Sr(Y_{0.5}ND_{0.5})O_3$ (see Fig. 6.3).

Sr(Ni_{0.5}Mo_{0.5})O₃, Sr(Co_{0.5}Mo_{0.5})O₃ and Sr(Zn_{0.5}Mo_{0.5})O₃ at temperatures of 300°, 400°, 430°, 230°, 320° and 420°C, reported for $Sr(Ni_{0.5}W_{0.5})O_3$, $Sr(Co_{0.5}W_{0.5})O_3$, $Sr(Zn_{0.5}W_{0.5})O_3$, Tetragonal to cubic phase transformations have been re-

spectively. The X-ray diffraction data indicates ordering of the B ions, but in Fig. 6.4 the cell size used for convenience was that of the simple perovskite structure. This figure shows the variation in cell size with temperature.

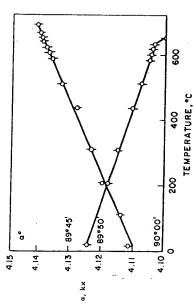


Fig. 6.3. Parameters of $Sr(Y_{0.6}Ta_{0.6})O_{8}$ as a function of temperature (after Smolenskii et al.(18)).

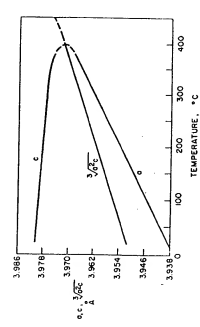


Fig. 6.4. Parameters of Sr(Co., Wo., 6)O, as a function of temperature (after Kupriyanov et al. (19))

Filip'ev and Fesenko studied the phase transition in $\mathrm{Pb}(\mathrm{Co}_{0.5}\mathrm{W}_{0.5})\mathrm{O}_{3}$ and found that it involved a change in symmetry from orthorhombic to cubic at 25°C. Superstructure was observed in both phases. A peak in the dielectric con-

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stant was seen at the phase transition, indicating that the compound was ferro- or antiferroelectric.

atures at which the peak in the dielectric constants occurred Antiferroelectric phase transitions from monoclinic to cubic symmetry were reported for the compounds $\mathrm{Pb}(\mathrm{Lu}_{0.5}\mathrm{Nb}_{0.5})\mathrm{O_3},$ Pb(Luo,Tao,s)O3 and Pb(Ybo,sNbo,s)O3 at 280°C. However, the transition temperature does not coincide with the temperfor these compounds.

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CHAPTER 7

FERROMAGNETISM

THE common exchange energy in magnetic oxides is of the indirect (super-exchange) type. The energy between spins of neighboring metal ions in perovskite-type structures is often found to be negative so that antiparallel alignment has the lowest energy. It has been proposed that this alignment is caused by mutual interaction of the metal ions with the oxygen ion which is situated between them.

The magnetic oxides with the perovskite structure,

found for compounds containing Co3+ and Co4+ ions, but not tween Mn3+ ions, a negative interaction between Mn4+ ions and strong positive interaction between Mn3+ and Mn4+ ions the moments of Mn3+ and Mn4+ ions and the highest values crystals of (La, Ca)MnO₃. A corresponding situation was or compounds containing Cr3+ and Cr4+, or with Fe3+ and La3+Mn3+O3-Ca2+Mn4+O3 and La3+Co3+O3-Sr2+Co4+O3, studied by Jonker and van Santen appear to be exceptions. (1) These studies indicated that a weak magnetic interaction beexisted in the manganites. It was found that in the solid soluhion range LaMn²⁺O₃ - 25-35% AMn⁴⁺O₃ (A = Ca, Sr, or Ba) the magnetic saturation values agreed with the sum of of the Curie temperatures occurred in this region. The saturation magnetization at 90°K is given in Fig. 7.1 for mixed All of these phases with the B metal ions in two valence states Fe⁴⁺, as the B ions were found to be antiferromagnetic⁽²⁾. $La^{3+}Mn^{3+}O_{3}-Ba^{2+}Mn^{4+}O_{3}, La^{3+}Mn^{3+}O_{3}-Sr^{2+}Mn^{4+}O_{3},$ were highly conducting.

In order to avoid the high conductivity, Jonker studied $\operatorname{La}(\operatorname{Mn}_{1-x}^{3+}\operatorname{Cr}_x^{3+})\operatorname{O}_3$ solid solutions which contained B ions of different elements with the same electronic configuration as the Mn^{3+} and Mn^{4+} ions. Positive magnetic interactions were

found between Mn³+-Mn³+ and Mn³+-Cr³+ pairs at low temperatures. The saturation magnetization increased up to a composition of 30% LaCrO₃ and then the increasing number of Cr³+-Cr³+ strong negative interactions lowered the saturation magnetization. The maximum was found in the region of the transition point from a monoclinic to a pseudocubic structure.⁽³⁾

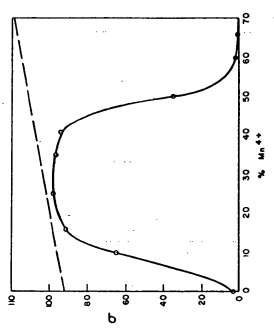


Fig. 7.1. Saturation magnetization at 20.4°K of LaMnO₃-CaMnO₃ (efter G. H. Jonker, *Physica* 22, 707 (1956)).

In a similar study of the (La,Ba)(Mn³+,Ti⁴+)O₃ system, Jonker found that a maximum in the saturation magnetization existed at the composition which produced a change from a monoclinic to a cubic structure. Since the titanum ion is diamagnetic the positive interaction between Mn³+ ions is the only one possible in these phases.

Goodenough et $al.^{(4.5)}$ investigated the $La(Mn_{1-x}Co_x)O_3$ series of phases and reported that the ferromagnetic saturation varied nearly linearly between x=0.20 and x=0.70. This was attributed to a positive interaction between Mn ions, with Co ions being in the diamagnetic low spin state. However, the similarity between the ferromagnetic satura-

be another explanation for the magnetic behavior of the forstrong positive interaction between the Co and Mn ions as boring pairs of Co and Mn ions were present in the divalent $\operatorname{La}(\operatorname{Mn}_{1-x}\operatorname{Co}_x)O_3$ series led Jonker to believe that there might mer series. (0) His studies indicated that there was a rather well as the positive interaction between Mn3+ ions suggested by Goodenough. In addition, the results suggested that neightion versus composition curve for the $La(Mn_1^{3+}Cr_x^{3+})O_3$ and

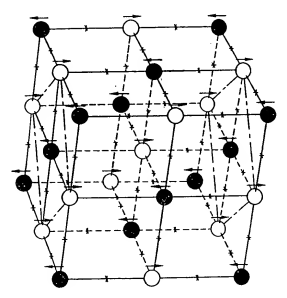


Fig. 7.2. Magnetic ordering in $Ba(B_{6,6}'B_{6,6}')O_3$ type compounds (after Longo and $Ward^{(6)}$).

and tetravalent valence states respectively in the solid solutions.

Wold $et al.^{(n)}$ reported ferromagnetism in the La $(Mn_{1-x}Ni_x)O_3$ system. This may be caused by Ni3+-Mn3+ and Ni3+-Ni3+ interactions. However, it is possible that the $\mathrm{Mn^{4}}^{+}$ and $\mathrm{Ni^{2}}^{+}$ may be present in some sites and also could interact.

Sugawara and Lida(8) prepared BiMnO₃ at 40 kbar and 700°C. The compound was found to be ferromagnetic at 130°K.

of 43°C, 128°C and 265°C, respectively. By analogy with the and Ca(Fe_{0.5}Re_{0.5})O₃ were found to have Curie temperatures Fig. 7.2. The compounds Ba(Fe_{0.5}Re_{0.5})O₃, Sr(Fe_{0.5}Re_{0.5})O₃ the general formula $A(B_0', \bar{b}_0'', O_3$ where \bar{B}' is \bar{W}^{5+} , Mo^{5+} , or that a negative interaction between the B' and B" exists.(9) The ordered arrangement of the two B cations is shown in In the magnetic-ordered perovskite-type compounds with Re* and B" also is a parametric ion, Longo and Ward propose

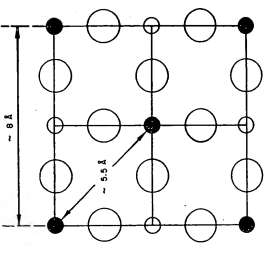


Fig. 7.3. Face of cubic ordered perovskite structure (100). $\bullet = B', O = B'', O = 0, A$ ions omitted (after Blasse⁽¹³⁾)

found for Sr(Fe_{0.5}Mo_{9.5})O₃ at an Fe-O-Mo bond distance of $A(Fe_{0.5}Mo_{0.5})O_{3}$ compounds showed a lower Curie temperature (Ba, Sr, Ca)(Feo.s Moo.s)O3 a maximum in the Curie point was magnetic properties of these compounds, an increase in Curie anticipated for compounds containing W5+ and Mo5+ as the B" ion. However, the $A(Cr_{0.5}Mo_{0.5})O_3$, $A(Cr_{0.5}Mo_{0.5})O_3$ and for Ca2+ compounds. (10) In fact, for the phases in the system temperature with decreasing A cation size might have been approximately 3.95 Å.(11) FERROMAGNETISM

nealing the compounds, which increased the ordering of the B ions and decreased the number of antiferromagnetic B'-0-Table 7.1 presents the ferromagnetic perovskite compounds Mg, Co, Ni and Cu), found that the magnetic exchange interactions between B' and Mn4+ ions in ferromagnetic compounds appear to be positive. (12) The saturation moment of $\mathrm{La}(\mathrm{Co_{0.5}Mn_{0.5}})\mathrm{O_3}$ and $\mathrm{La}(\mathrm{Ni_{0.5}Mn_{0.5}})\mathrm{O_3}$ was increased by an-B' and Mn*+-O-Mn*+ interactions, so that a decreasing Curie temperature was expected and found experimentally. Blasse, in a study of perovskites $La(B_{0.5}'Mn_{0.5})O_3$ (B' = and their Curie points.

TABLE 7.1. Ferromagnetic Compounds

	Curie temp.	References
Barremood	64	10, 11
Srremood	146	10, 11
Carremood	104	10
Srchwood	200	10
Carchwood	- 125	10
Srchwood	180	10
Sarremood	- 130	10
Barremood	43	9
Srremood	43	9
Srremood	43	9
Srremood	128	9
Srremood	265	9
Carremood	Magnetic R.T.	17
Carremood	Magnetic R.T.	17

A(B',5B',5)O3 where B' is the only paramagnetic ion, Blasse showed that the mechanism for magnetic interaction between the paramagnetic B' ions is of the type B'-O-B''-O-B''(13) (see Fig. 7.3). This is a new type of superexchange in the perova study of antiferromagnetic ordered perovskites skite structure.

In a study of $Sr(B_{0,s}'W_{0,s})O_3$ type compounds where B' = Mn, Fe; Co and Ni, Blasse⁽¹⁴⁾ found that the Néel temperature (see Fig. 7.4) increased as it did in the B'O compounds

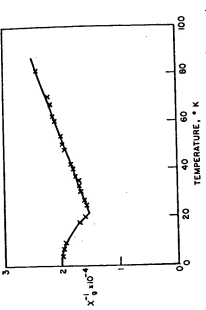


Fig. 7.4. Reciprocal susceptibility per gram vs. temperaturo of $Sr(Co_{0.6}W_{0.6})O_s$ (after Blasse⁽¹⁴⁾).

(rocksalt structure) and KB'F3 compounds (perovskite structure). See Table 7.2.

Table 7.2. Magnetic Properties of Compounds Sr(B_{0,5}W_{0,5})O₃, B'O and KB'F₃ (after Blasse⁽¹⁴⁾)

$KB'F_3$ T_N (°K)	88 113 114 275 243
$T_{ m N}$ (°K)	116 186 292 523
$\frac{\mathrm{Sr}(\mathrm{B_{6.8}W_{6.8}})\mathrm{O_3}}{T_{\mathrm{N}}}$	10 16 22 54 1
Bí	Mn ²⁺ Fe ²⁺ Co ²⁺ Ni ²⁺

Since the Néel temperature is linearly related to the exchange constant of the 180° B'-O-W-O-B' and B'-O-B' interaction is about 10 times as strong as the B'-0-W-0-B' actions, the data in Table 7.2 shows that the B'-0-B' inter-In the complex perovskite structures the B' ions are separated by an OWO array, and in MeO by a single oxygen ion. interaction.

In addition, from measurements on Ca(Fe_{0.5}Sb_{0.5})O₃ and

Sr(Fe_{0.5}Sb_{0.5})O₃, Blasse found that the magnetic interactions respectively. If incomplete ordering exists, such as is found in Sr(Mn_{0.5}Sb_{0.5})O₃, a strongly positive Mn³⁺-O-Mn³⁺ interbecame stronger for shorter distances, $T_{N}({}^{\circ}\mathbf{K}) = 31$ and 21, action can be present.

sitic ferromagnetism by careful control of sample stoichioskites. It was reported in GdFeO3, for temperatures between 78° and 295°K, (15) the magnetization varied for high fields where the parasitic ferromagnetization σ^0 amounted to about Wold (10) has prepared samples of LaFeO₃ with reduced parations should be a requirement before measurements are made 1% of the $\sigma(\text{Fe})$ available. It was attributed to imperfectly compensated antiferromagnetism of the Fe³⁺ ion sublattice. metry, concluding that the use of careful preparation condi-Parasitic ferromagnetism has also been observed in perov-(H > 6000 oe) according to the expression $\sigma = \sigma^0 + \chi H$, on ferromagnetic materials.

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CHAPTER

OPTICAL PROPERTIES

8.1. TRANSMITTANCE

of the crystal was nearly a constant value of $\sim 2.4 \, \, \mathrm{from} \, \, 20^\circ$ to 90°C and reached a maximum value of 2.46 at 120°C (see Fig. 8.1). At the transition temperature a sudden change tals of BaTiO₃ at various temperatures. The refractive index Merz(1) studied the optical properties of single-domain crysin n is observed (see Fig. 8.2).

Lawless and DeVries⁽²⁾ also measured the index of refracstant index of 2.368 was obtained from 20° to 105°C and tion of BaTiO3 at 5893 A in the range of 20-160°. A conabove the Curie point the index increased 1.3% to 2.398 and remained constant to 160°C.

The single crystal of BaTiO₃, 0.25 mm thick, was found to transmit from 0.5 μ to 6 μ . Complete absorption was found

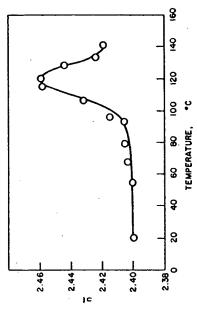


Fig. 8.1. Refractive index \vec{n} as a function of temperature (after Merz⁽¹⁾).

OFFICAL PROPERTIES

for wavelengths greater than 11 μ and a feeble absorption

band existed near 8 μ (see Fig. 8.3). The optical properties of strontlum titanate single crystals produced by a flame fusion process were reported by Noland. (3) The optical coefficient was obtained from 0.20 μ to 17 μ in wavelength (see Fig. 8.4). Transmission of better than 70% was measured from 0.55 μ to 5 μ . The index of

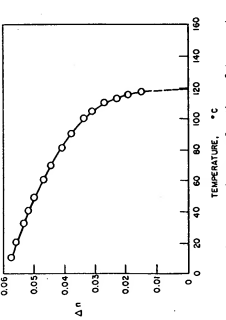
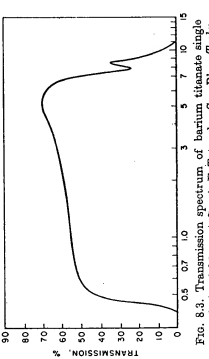


Fig. 8.2. Birefringence An as a function of temperature (after Merz⁽¹⁾).



crystal, d=0.25 mm (after A. F. Tatsenko, Sov. Phys. Techn. Phys. 8, 2257 (1957)).

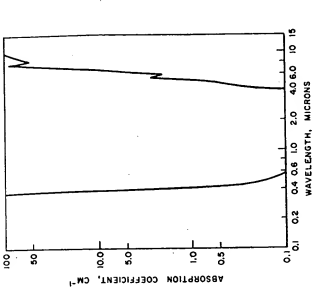
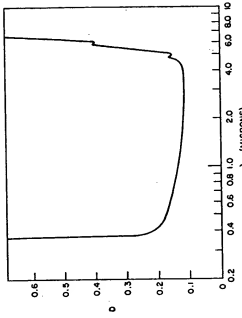


Fig. 8.4. Absorption coefficient of single crystal SrTiO₃ as a function of wavelength (after Noland⁽⁹⁾).



Fra. 8.5. Optical density vs. wavelength of 1.9 mm CaTiO₃ crystal (after Linz et al.(4)). X (MICRONS)

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refraction of these crystals is 2.407 at 5893 A, the dielectric constant is 310 and the loss tangent 0.00025.

technique (see Fig. 8.5). The absorption characteristics are quite similar to those of SrTiO3 crystals with the exception that the absorptions are shifted to shorter wavelengths. Index of refraction data for CaTiO3 and SrTiO3 as a function of wave-The optical density of CaTiOs was reported by Linz and Herrington. (4) The crystals were prepared by the flame fusion

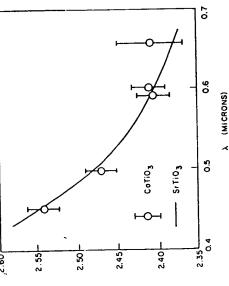


Fig. 8.6. Dispersion of CaTiO₃ and SrTiO₃ (after Linz et $aL^{(4)}$).

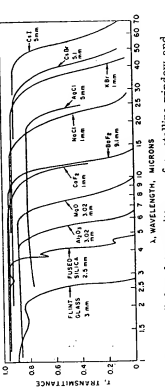


Fig. 8.7. Infrared transmittance of crystalline window and prism materials after L. J. Neuringer (Electrical Manufacturing) (March 1960), copyright (Conover-Mast publications,

length are plotted in Fig. 8.6. The data are quite similar and the deviation at 657 Å can be attributed to the difficulty

in the measuring technique at that wavelength.

being used are shown in Fig. 8.7. Note that the titanates are useful in the same wavelength range as silica and $\mathrm{Al_2O_3}$. In addition, strontium titanate is considered as an excellent For many applications the detector-lens combinations are cooled to solid CO, and liquid N, temperatures to increase the sensitivity. Salzberg(5) made successive transmittance measurements on strontium titanate from room temperature to -187°C showing that there was no decrease in the transmaterial for use with optically immersed infrared detectors. ture infrared windows. Typical materials which are presently BaTiO3 and SrTiO3 have been considered for high-temperamittance of strontium titanate down to -187°C.

8.2. Coloration by Light

heating reversed the effect. The effect presumably was caused nates took on a purplish color when exposed to light and MacNevin and Ogle⁽⁶⁾ noticed that the alkaline earth titaby impurities.

8.3. Electro-optic Effect

field of 13,000 V/cm, which corresponds to a retardation of 180 half-waves per cm of light path length at 6828 Å. Thus, for polarization of amplitude modulators, the power and voltage requirements can be satisfied by transistor circuitry. The large value of arrho n also should make possible a light-scanmeasured by Geusic et al." The electro-optic coefficients of ndicatrix are expressed in terms of the induced polarization.(7) Thus, the coefficients might be fundamental properties of the perovskite lattice. These studies also showed that KTN has a large room-temperature electro-optic effect, low electrical losses and a large saturation polarization. An induced birefringence of 0.0057 has been reported with an applied The electro-optic properties of KTaO₃, K(Ta_{0.65}Nb_{0.35})O₃ KTIN), BaTiO3 and SrTiO3 in the paraelectric phase were these perovskites are nearly constant with temperature and from material to material when the distortions of the optical

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ning system using a prism or a partially electroded cube which can scan over 500 resolvable beam diameters.

Cohen and Gordon⁽⁸⁾ described experiments which involve propagating an electromagnetic wave through an electrooptic medium. Using 2.5 µsec microwave pulses with a peak power of a few hundred watts at 9.5 GC, 5% of the light in the zero-order beam was transferred to the higher orders at frequencies 750 GC. Therefore, when the beam is polarized at 45° to the direction of the microwave electric field, it should be possible to extinguish the zero-order beam and pass 75% of the deflected beam.

I. P. Kaminow⁽⁹⁾ described experiments in which ferro-

I.P. Kaminow⁽⁹⁾ described experiments in which ferroelectric barium titanate was used at 70 MC as an optical phase modulator. Focused fundamental gaussian mode passed through the edge of a crystal plate with both the higher order "donut" mode of a 0.633- μ He–Ne maser and the fundamental mode maintaining their identity.

in addition, the materials with the higher dielectric constants from 50 to 500 Å (less than the wavelength of light) and the tallite sizes had the higher dielectric constants. In these as a function of electric field. No electro-optical activity was containing small amounts of cadmium were formed by conelectric NaNbO3 ferroelectric. The crystal size was varied dielectric constant from 50 for the glass to 550 for the transparent crystallized material. Materials which had larger orysexperiments the difference in refractive index was measured observed for the glass, but the retardation was proportional to the square of the electric field for the crystallized glass. Borrelli et al. (10) observed the electro-optic effect of ferroelectric microcrystals in a glass matrix. Crystals of $\mathrm{NaNbO_3}$ trolled crystallization of a glass with a high silica glass remaining as a matrix. The cadmium was used to make antiferroproduce the largest index differences, see Fig. 8.8.

8.4. LASERS

In recent years there has been considerable interest in materials to be used for laser application. The operating laser systems are listed in Table 8.1.

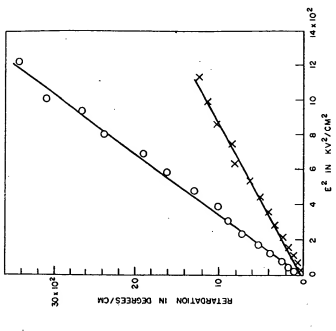


Fig. 8.8. Retardation in degrees of arc per om vs. the square of the applied electric field (O, sample with ε of 540; \times , sample with ε of 356). Measurements made at 6328 Å (after Borrelli⁽¹⁰⁾).

In selecting perovskite laser host materials a great deal can be learned from an examination of these systems.

The ion Nd³+ appears to be the most popular for introduction into relatively large crystallographic sites. However, except when LaF₃ is used as a host, compensating ions are required in these substitutions. Divalent Tm²+ and Dy²+ can be substituted in CaF₂ without compensating ions but these divalent rare earth ions are relatively unstable. For Al₂O₃ whose structure contains octahedrally coordinated crystallographic sites for the Al³+, Cr³+ proved to be ideal for substitution. Other small ions which have been used to produce laser action are Fe³+ and Mn⁴+, but these ions have not been employed to the extent Cr³+ has.

The perovskite structure should be a promising one since

Table 8.1. Data for Operating Laser Systems

Tagar	(#)	temp. (°K)
1. Al ₂ O ₈ (Cr ³⁺)	0.6944	300 (Cont.)
2. CaF ₂ (U ³⁺)	2.556	77 (Cont.)
	2.407	11
BaF,	2.556	77
٠.	1.06	300
Ca.W(1.0646	300 (Cont.)
7. SrMoO (Nd8+)	1.0643	
	1.064	77 (Cont.)
8. CaMoO (Nd3+)	1.067	300
 PbMoO₄(Nd³+) 	1.0586	300
10. NaLaMoO.		
(Nd3+)	1.0586	300
1. CaF,(Nd3+)	1.046	300
12. SrF,(Nd3+)	1.06	300
3. BaF ₂ (Nd ³⁺)	1.06	300
14. LaF ₃ (Nd ³⁺)	1.06	300
15. CaWO4(Ho3+)	2.046	11
 CaF₂(Ho³⁺) 	ю	11
Ξ.	1.116	7.7
٠.	1.91	300
. Ca.WO.	1.612	11
20. CaWO (Pr ³⁺)	1.05	11
21. Glass (Yb3+)	1.01	7.7
22. Glass (Sm³+)	0.71	77
23. Glass (Gd3+)	0.31	
₩.		300 (Cont.)
		0,000
	-	300 (Cont.)
-	0.7082	: '
27. $CaF_2(Tm^{2+})$	1.116	4 (Cont.)
	6.9	

ions. However, as was pointed out previously no simple cubic perovskite-type compounds exist which will accept trivalent activator ions without producing defects or require charge it contains a large A site suitable for Nd3+ and smaller B site for Cr3+. In the ideal structure these sites are cubic and centsary for obtaining long fluorescence lifetimes for activator rosymmetric, a condition theoreticians feel should be neces-

crystals big enough for fluorescence measurements have been prepared. (12) The measurements showed the fluorescence line material. However, flux-grown and hydrothermally grown metry. The fluorescence lifetime for Cr3+ in LaAlO, has been reported to be as high as 46 msec as compared with 3 msec time has been attributed to the nearly cubic symmetry of the site containing the Cr3+ ion. The transition of LaAlOs at 435°C presents a problem in obtaining single crystals of thin compensating ions. The compound LaAlO₃, however, has a structure which is only slightly distorted from cubic symfor the lifetime of Cr3+ in Al₂O₃. (11) This enhancement in lifeat 7356 Å which is characteristic of Cr3+.

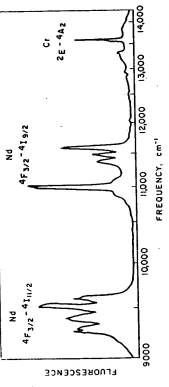


Fig. 8.9. Fluorescence spectrum of LaAlO₃-Cr, Nd excited with high-pressure mercury lamp (after Murphy et $aL^{(11)}$).

studies indicated that energy transfer existed between the $^4\mathrm{T}_2$ level of Cr^{3+} and the D levels of Nd^{3+} . the remaining two groups are associated with the fluorescence of Nd3+. Lifetimes of Md3+ and Cr3+ were found to be 0.46 der samples. The fluorescence spectrum of LaAlO₃: Cr, Nd is presented in Fig. 8.9. The line at 7356 Å is due to Cr3+ and and 4.6 msec, respectively, at room temperature. Their Murphy et al. $^{(11)}$ reported on energy transfer between Cr^{3+} and Nd3+ in LaAlO3. These studies were conducted on pow-

of the studies was to increase the efficiency of lasers from the radiation from Cr3+ to Nd3+ ions in GdAlO3. The purpose usual 2% or less. The low efficiency was attributed to the poor Ohlmann and Mazelsky⁽¹³⁾ also found transfer of excitation

spectral mismatch between the broad bands of the pump lamps and the relatively narrow absorption bands of the paramagnetic ions. Thus, Cr3+ was added to GdAlO3 to produce broad absorption bands which could transfer the energy it absorbed to the Nd³+ ion. The fluorescence spectrum showed fluorescence at 0.727 μ due to chromium. When the Cr³⁺ evels were excited, the fluorescence of neodymium showed an ples over the singly doped samples. The lifetimes of Md³+ and two groups occurring around 0.9 μ and 1.07 μ from Nd $^{3+}$ and intensity increase of several times in the doubly doped sam-Cr3+ in GdAlO₃ were 0.130 msec and 15 msec, respectively.

Other simple perovskite compounds considered for laser However, divalent or tetravalent laser activating ions would be required for substitution so that the symmetry would not be destroyed by compensating ions. The activating ions in host materials were SrTiO3, SrSnO3, BaSnO3 and BaZrO3. these valence states are not particularly stable.

indicate that there may be some enhancement in lifetimes due to cubic symmetry about the substituted ion. However, the increase is small and it appears that the neodymium lifetime selected since compounds of this type containing barium are usually least distorted, and Ta5+ is one of the B ions because of its resistance to reduction and the other B ion should not produce energy levels which would interfere with those of the doping ions. The fluorescence lifetimes of powders of these compounds containing activator ions are presented in Table 8.2. The values obtained, i.e. 850 msec for Nd3+ in $\mathrm{Ba(Lu_{0.5}Ta_{0.5})O_3}$ and 800 msec in $\mathrm{Nd^{3+}}$ in $\mathrm{Ba(Gd_{0.5}Ta_{0.5})O_3},$ ions. They found that compounds of the Ba(B3+Tao.s)O3 had an ordered structure in which the B ions alternate so that the symmetry about the B site is retained. Barium was Because of the enhancement of lifetimes observed for Cr3+ n LaAlO3 and GdAlO3 as compared with its lifetime in Al₂O₃, a search was made by Galasso et al. (14) for cubic perovskite compounds which would accept trivalent rare earth s not influenced as much by environment as that of Cr3+.

Ba(Gdo.,Nbo.,s)O3 host confirmed the fact that a center of symmetry existed at the B site in the structure of these cubic The fluorescence spectra of Er3+ ion studied in complex oxides.(15)

Table 8.2. Fluorescent Lifetime Data (after Galasso et al. 41)

Doped Ba(B _{0.8} Ta _{0.8})O _{\$} phases Nd*+ doped	Emission line (µ)	Lifetime at 300°K (µ sec)	Lifetime at 77°K (µsec)
Ba(La _{0, 48} Nd _{0,02} Ta _{0,50})O ₃ Ba(Gd _{0,48} Nd _{0,02} Ta _{0,50})O ₃ Ba(X _{0,48} Nd _{0,02} Ta _{0,50})O ₈ Ba(Lu _{0,48} Nd _{0,02} Ta _{0,50})O ₈ Ba(Lu _{0,48} Nd _{0,02} Ta _{0,50})O ₈ Ba(Ru _{0,48} Nd _{0,02} Ta _{0,50})O ₈ Ba(Sc _{0,48} Nd _{0,02} Ta _{0,50})O ₈	1.06	200 200 400 150 200	650 800 250 850 250 220
$\mathrm{Sm^{3+}}$ doped $\mathrm{Ba}(\mathrm{Y_{0.48}Sm_{0.02}Ta_{0.60}})\mathrm{O_{3}}$	0.70	1450	
$ m Yb^{3+}$ doped $ m Ba(Y_{0.48}Yb_{0.02}Ta_{0.80})O_{8}$	1.01	1800	!

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CHAPTER 9

OTHER PROPERTIES

9.1. CATALYSTS

Fuel Cell

A detailed study was conducted by Epperly et al. to evaluate complex perovskites of the general formula $A(B_0',B_0',S)O_3$ with A = Li, Cs, Ca, Sr, Ba, La, Tl, Pb and Bi, B' = V, Cr, Mn, Fe, Co, Ni, Ru, Rh, Ir and Pt and B'' = Ti, Zr, Sn, Hf, V, Nb, Ta, Mo and W for fuel cell application. The B' ions were selected from those which are known to have catalytic properties and B'' ions were selected for the corresion resistance they impart to a compound. The problem was to tailor the compounds so that they would have high conductivity and high resistance to attack by dilute solutions of sulfuric acid. While it is dangerous to generalize, the information generated can be quite useful if it is realized that it is qualitative in nature.

They found that most of the perovskite compounds studied were quite acid resistant, but were poor conductors. It is important to note that the B ions were used in their highest oxidation states. The incorporation of oxygen vacancies improved the conductivity but the acid resistance was not as good. Compounds containing Ni had good acid resistance, especially when Ni²⁺ ions were added to the acid, but were poor conductors. The compounds containing cobalt were made conductive by producing oxygen vacancies, but then became soluble in the acid. Compounds containing Nb or Ta were more acid resistant than those containing Mo or W. Compounds containing Ti, Zr or Hf had relatively poor acid containing manganese in more than one valence state.

The tungsten bronzes also have been considered for elec-

trodes in fuel cells. Dickens and Whittingham⁽²⁾ studied the recombination of oxygen atoms on the surfaces of phases with the formula M_xWO_3 where M is Li, Na or K and 0 < x < 0.80. The pattern of catalytic activity for oxygen atom recombination is approximately the same for Li_xWO₃, Na_xWO₃ and K_xWO₃ phases. In the semiconducting range, from x = 0 to x = 0.25, the catalytic activity decreased, but in the range above x = 0.25 the catalytic activity increased sharply to a maximum at x = 0.55. Above x = 0.25 the conductivity of the bronzes is metallic. Thus, the catalytic activities of the tungsten bronzes appear to be closely related to the electronic properties of the bronzes.

Heterogeneous Catalysis

Parravano⁽³⁾ studied the effect of the electronic rearrangement of ferroelectric materials at their transition temperatures on the electron exchange occurring during a catalytic process. From data obtained for carbon monoxide oxidation on KNbO₃, he concluded that the change in the rate of catalytic oxidation of CO is affected by the electronic transition. This supported the evidence for an electronic mechanism as being rate determining for the catalytic oxidation of CO.

9.2. THERMAL CONDUCTIVITY

The thermal conductivity values for the titanates are listed in Table 9.1. They are quite low, with the conductivity of barium titanate being lower than that of strontium titanate and calcium titanate at 50°C.

9.3. Melting Points

The melting points of many of the ternary oxides have been determined, but none of complex perovskite type compounds (see Table 9.1). It should be realized that in many cases these are only approximate values; however, they do serve as guides for crystal growth experiments, for example, note that zirconates are extremely high melting-point materials along with hafnates and thorates, the titanates have

Phase	Thermal conductivity (W/cm°C)	Coeff. of thermal Expansion (×10-6/°C)	/kcal/mole)	Melting point (°C)
BaThO, BaTiO,	0.022-0.032	19 (10–90)	394.8	2299 1610
BaZrO ₃	(90-230)	97.		2688
CaHfO3		25		2471
CaTiO	0.043-0.046	7.08 (1000-1300) $13.04 (25-500)$	397.4	1975
CaZrO	(90-139)			2343
GdAlO3 KNbO3				1039
KTaU3 LaAlO3				2075- 2080
LaFeOs				1888
NaN bO3 NaTaO3				1780
PbTiO _s		-16 (below 490) 25 (above 490°)		
SrTiO ₃	0.037-0.058	8.63	398.9	2080
SrZrO3	(0#1-00)	8.84 (25–500)		2799
YAIO.		_		1950
Ba(Fe, Ta, b)		10.99		
			_	

intermediate melting points, and the niobates and tantalates are relatively low melting.

9.4. HEATS OF FORMATION

measured. These values shown in Table 9.1 are quite similar Only the heats of formation of the titanates have been for barium, strontium and calcium titanate.

9.5. THERMAL EXPANSION

BaZrO3, while the coefficient of thermal expansion of titanate is unusual, having a negative coefficient up to 490°C of thermal expansion of barium titanate above the Curie point is similar to that of calcium titanate and larger than that of strontium titanate. For the zirconates the coefficient of thermal expansion of SrZrO₃ is greater than that for Ba(Fe, Ta, Ta, S)O3 is greater than that of Sr(Fe, Ta, S)O3. Lead sion of the perovskites in relation to the ion in the A position (see Table 9.1). Note that for the titanates, the coefficient and above this temperature it behaves like the other tita-There does not appear to be a trend in the thermal expan-

9.6. Density

The densities of perovskite-type compounds were calculated from X-ray data using the equation in a computer program

 $D = \frac{1}{\text{Volume of unit cell} \times 6.023 \times 10^{23} \text{ molecules/mole}}$ Molecular weight Xno. of molecules per unit cell

In the program, the equation for the volume of a triclinic cell was used $V = a \times b \times c / (1 - \cos^2 \alpha - \cos^2 \beta - \cos^2 \gamma + 2 \cos \alpha \cos \beta \cos \gamma)$

ed by introducing the periodic table with atomic weights the appropriate atomic symbols and multipliers. The cube duced for each compound. The molecular weight was obtaininto the computer and then inserting the data deck with and the appropriate parameters from Table 2.2 were intro-

no. of molecules per unit cell volume of the unit cell

was also calculated to obtain a', the unit cell edge of a cubic cell which represents the same volume allowed in the true unit cell for a molecule of a perovskite-type compound. This permits a comparison to be made between the ionic radii of the ions in these compounds even when they are indexed on cells with different symmetry. Table 9.2 presents the molecular weight, a', and the density of the perovskite compounds.

9.7. MECHANICAL PROPERTIES

cal properties of perovskites (see Table 9.1). The modulus Only very little information is available on the mechaniof barium titanate has been measured to be 16×10^6 psi.

the bend strength 13×10^3 psi and the compressive strength strain curve as shown in Fig. 9.1. (4) The small yield observed was attributed to domain motion or crystal twinning similar to that which takes place in ferroelectric BaTiO3 under The modulus of SrZrO₃ has been found to be 12×10° psi, 5×10^3 psi. Strontium zirconate has an interesting stressan electric stress.

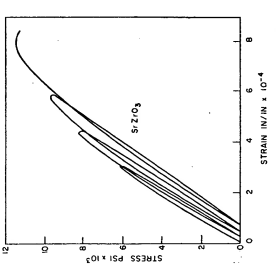


Fig. 9.1. Stress-strain behavior of hot-pressed SrZrO₃ (after Tinklepaugh⁽⁴⁾).

OTHER PROPERTIES

Volume

Formula

Compound	weight	Volume	Z	a', A	Density
	A1+E	A1+B6+O3			
AgNbO ₃	248.774	974.325	16	3.934	6.782
AgTaO ₃	336.816	60.481	-	3.926	9.244
CsIO	307.808	101.195	_	4.660	6.049
KIOJ	214.005	85.752	7	4.410	4.143
KNbO ₃	180.006	129.368	67	4.014	4.620
KTeO,	268.048	63.450	-	3.988	7.013
NaNbo,	163.894	119.427	87	3.808	4.556
NaTaO	251.936	117.498	67	3.887	7.119
RbIO,	260.373	92.345	7	4.520	4.680
T'IIO,"	379.273	91.716		4.510	6.865
	<u>1</u> +• •	(
	Τ. - ₩	A-1 D-1 O3			
BaCeO.	325.458	85.010		4.397	6.355
BaFeO,	241.185	63.520	_	3.990	6.303
BaMoO,	281.278	65.929	~	4.040	7.079
$\mathbf{BaPbO_{\tilde{s}}}$	392.528	78.019	-	4.273	8.352
BaPrO,	326.245	82.540	-	4.354	6.561
$B_{\mathbf{a}}P_{\mathbf{u}}O_{\mathbf{a}}$	427.338	84.605	_	4.380	8.385
BaSnO,	304.028	69.782	_	4.117	7.232
BaThO	417.376	90.217	7	4.485	7.680
BaTiO,	233.238	64.110	~	4.002	6.039
BaUO	423.368	84.431	–	4.387	8.324
$BaZrO_s$	276.558	73.665	_	4.192	6.232
CaCeO,	228.198	460.683	∞	3.862	6.578
CaHfO,	266.568	1	1 9	1	6
$CaMnO_s$	143.016	833.656	20 (4.706	2.270
$CaMoO_3$	184.018	472.590	00	3.896	0.171
CaSnO	206.768	246.406	4	3.949	5.672
$CaThO_3$	320.116	667.628	œ	4.370	6.368
CaTiO	135.978	223.913	7	3.825	4.032
CaUO	326.108	286.060	4	4.161	7.570
CaVO	139.020	216.125	4	3.775	4.291
CaZrO ₃	179.298	257.617	4	4.008	4.621
CdCeO,	300.518	447.697	œ	3.825	8.914
CdSnO	279.088	243.371	4	3.933	7.616
CdThO,	392.436	667.628	œ	4.370	7.806
CdTiO	208.298	218.491	4	3.794	6.330
CdZrO,	251.618	1	Ī	1	1
EuTiO	47	69.182	_	3.897	6.952
$MgCeO_s$	212.430	622.836	∞	4.270	4.529
	-				

Density

α', A

N

Volume

Formula weight

Table 9.2 (cont.)

A*+B*+O* (cont.)

6.638 6.532 7.300 6.773 6.631 7.208 4.468 7.212 7.212 7.212 6.441 6.446 6.441 6.446 6.441

3.870 3.887 3.788 3.824 3.883 3.895 4.110 3.896 3.838 3.940 3.838 3.940 3.838

250.297 229.560 64.354 65.906 234.202 243.086 236.413 277.773 462.246 61.163 266.089

411444446141

TABLE 9.2 (cont.)

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Compound		Control	GdVO ₃	LaAlOs	LacoC	LaCrO _s	To Go	To India	CiV.	LaBhO.		I.a.T.i.O.		T CO		Contraction	Nacoo.	NAMPO.	C COLL	Nagae Nagao	Notice of the second	Namos Nago	Masses 3	Pr A 10	CoOrd	PrOrO.	Pr.FeO.	PrGeO.	PrMnO	PrScO.	PrVO.	PuAIO,	PuCro,	PuMnO	P ₁ VO.	Sm A10.		SmCrO	S. T. N.	SmInO.	Sommer S	VAIO.	I Days
										òn -				•																									-				
Density			11.865	9.885	8.014	6.756	5.854	5.605	5.488	7.740	6.120	7.828	ı	6.434	7.068	6.120	7.780	5.458			8.201	8.715	8.430	6.632	868.9	6.827	7.333	ı	6.690	8.411	7.702	7.820	8.699	7.243	7.604	7.333	8.331	7.440	8.397	7.662	7.531	7.748	
á, À			3.810	3.974	3.974	4.640	4.276	3.862	3.869	4.069	3.975	4.174	l	4.033	4.420	3.904	4.304	4.101			3.859	3.890	3.946	3.776	3.866	3.900	3.879	1	3.900	3.936	3.713	3.838	3.700	3.733	3.803	3.869	3.965	3.728	3.738	3.820	3.861	3.820	
12	=		<u></u>	∞	-	00	4	œ	7	_	_	4	1	_	00	7	4	4	•		œ	, –	_	_		7	_	1	74	<u>∞</u>	4	4	7	4	_	7	7	4	-	4	4	-	_
Volume		s (cont.)	442.451	502.268	62.780	799.179	312.783	460.994	57.916	67.369	62.812	290.801	i	65.617	690.807	59.502	318,903	275.952	-	3+0 3	459 822	58.851	61.424	53.838	57.781	59.319	58.366	1	59.319	487.813	204.722	226.162	50.653	208.034	55.002	231.631	62.335	207.251	52.228	222.929	230.217	55.743	I
Formula	weight	A2+B4+O8 (cont.)	395.308	373.878	303.088	346.408	275.738	194.551	191.465	314.108	231.558	342.808	236.688	254.308	367.656	183.518	373.648	226.838	-	A3+B3+O,	983 960	308.974	311.916	215.100	240.114	243.965	257.838	233.074	239.060	308.974	237.480	266.345	265.436	226.940	251.954	255.805	312.825	232.230	264.181	257.244	261.095	260.186	
Dunoamon	nunodino		PbCeO ₃	PhsnO,	PbTiO,	PbZrO,	SrCeO,	SrCoO	SrFeO	SrHfO	SrMoO,	SrPbO,	SrRuO	SrSnO。	SrThO.	SrTiO,	Sruo	SrZrO	•		Bi A10	BiCrO	BiMnO	CeAlO	Cecro	CeFeO.	CeGaO	CeScO,	CeVO,	CrBiO,	DyAlO,	DyFeO,	DyMnO.	EuAlO,	EuCrO,	EuFeO	FeBiO.	GdAlO	GdCpO.	Gacco	GdFeO.	GdMnO.	

6.890 7.738 7.007 7.561 7.442 7.442 7.442 6.122 6.868 9.486 7.575 6.369 6.789 7.262 6.789 7.262 7.262 7.362

3.752 3.752 3.853 3.860 3.886 4.068 3.860 3.3787 3.811 3.811 3.811 3.811 3.811 3.811 3.811 3.810

> 244.752 258.625 243.843 233.861 239.847 316.980

228.799 235.092 230.064 269.200 64.872 257.276 235.115 37.778 64.310 230.181 239.385 232.103 65.743

62.819 63.882

60.236 63.521

256.204
256.190
213.890
245.841
238.904
242.755
256.628
186.908
2242.755
225.813
221.864
234.808
237.850
275.813
219.220
261.171
244.234
248.086
261.958
192.238
247.176
224.136
221.868

260.334 235.641 97.324 233.456 67.512 239.179 208.928

53.123 226.109 232.589 265.872 58.864 203.404

341.994 344.936 340.940 225.330 257.281 254.195 198.348 249.290 Density

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6.554 6.523 6.911

4.120 4.350 4.187

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OTHER PROPERTIES

PREPARATION OF PEROVSKITE-TYPE COMPOUNDS 148

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cont
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9.2
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TABLE 9.2 (cont.)						TABLE 9.2 (cont.)			
H F	Formula weight	Volume	Z	a', A	Density	Compound	Formula weight	Volume	
⋖	3+B3+(A3+B3+O, (cont.)					Δ/R' R"	A/B' B" 10 (cont	-1 ~
188	188.899	218.305	4	3.793	5.746		0-0.69-0.0		٠-
192	192.750	225.862	4	3.836	5.667	Ba(Sc. 67 Wo. 33) Os	821.97.2	009.470	
181.859	829	244.887	4	3.941	4.931	Ba(Yo.67,U0.33)O3	323.404	500.000	
	. ✓	A BO				D8 (X 0.67 W 0.33) O3	361.945	1 1	
	H	, 1				DL (12 117 10 38)	353 976	ļ	
187	187.144	119.550	~	3.910	5.197	2-(C, D) (1) (1) (1) (1) (1) (1)	931 909	513,922	
275	275.186	119.857	~	3.913	7.623	OF (OF 0.67 IVE 0.89) C3	200.00	000 613	
288	282.571	113.684	67	3.845	8.252	Sr(Cr _{0.67} C _{0.33})C ₃	248.000	012:000	
86	980 839	118 370	6	3.875	8.012	$Sr(Fe_{0.67}Re_{0.33})O_3$	234.482	491.109	
1 -	200:00	190 776	16	3 003	5 123	Sr(Fe _{0.6} , W _{0.33})O ₃	233.706	61.490	
9 5	# 6	7.00	9 0	2000	10.100	Sr(In, "Re, ")O,	197.064	571.167	_
7.7	274.786	121.087	.7	3.927	7.534	T. (Co. In D. In D.	257.052	245.666	_
180	188.503	118.332	87	3.897	5.289	1 2 C C C C C C C C C C C C C C C C C C	966 571	244 188	
27(276.545	118.789	83	3.902	7.729	La(Co.670 Do.33/O3	7.0.00	007:327	_
18,	187.404	119.092	83	3.905	5.224		A(B',	A(B' ₃₃ B'' ₆₇)O ₃	
97.6	975 446	119 246	S	3.907	7.669		010 000	300 000	
	010.540	21011	0	200	7 886	Ba(Cao, 33 N Do. 67) U3	200.012	0.00.040	
7	700.0	#17:11	4 (0000	000.7	Ba(Ca _{0,33} Ta _{0,67})O ₃	319.800	219.214	
726	268.286	113.227	2	3.840	7.573	Ba(Cd, "Nb, 17)0,	284.677	72.407	
28(286.049	110.852	N	3.813	8.567	Ba(CdTa)O.	343.665	72.355	
248	248.986	446.725	4	4.816	3.701	Be (Co. N.P.) O.	267.033	68.418	
23	238.787	51.479	7	3.720	7.700	D&(Co. 33-1 Co. 67) (3	396.091	204 817	
25	254.838	57.602	7	3.862	7.344	D&(CO. 33 L &0.67) O3	960.050	649 989	
06	909 938	63 099	-	3 983	5.321	Da(Cu _{0.88} N D _{0.67})O ₃	200.000	200.00	
3 6	007.700	125.00	٠,	40.6	5 745	Ba(Fe _{0.83} Nb _{0.67})O ₃	200.010	00.00	
7 -	7 F	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	•	010.#	-	Ba(Fe _{0.33} Ta _{0.67})O ₃	325.003	68.921	
4 ,	143.000	l	1	l	•	Ba(Mg _{0.83} Nh _{0.67})O ₈	255.608	204.134	
6 	194.535	I	I	1	1	Ba(Mg, "Ta, ")O,	314.596	204.608	
19	191.449		I		1	Ba(Mn, 2, Nb, 2, 1)O.	265.715	1	
18	183.502	İ	I	1	ı	Be (Mrs. Tra. 1)	324,703	208.994	
<u>8</u>	186.544]	I	1	1	Ba(Ni,Nb,)O.	266.960	67.618	
	į					Bo (Ni The)O	325 948	202.482	
	A(B).83	A(E _{0.67} E _{0.33})U ₃				Bo (Dh. 101)	315.958	77.309	
26	264.086		Ī	ļ	!	Bo (D) To)O	374.946	76.768	
33	354.884	589.745	00	4.193	7.991	De(1.00.33.140.67)(3	325 489	929.026	
8	358 073	589 745	α	4.193	8.063	D&(OLO.33 L &0.67) O3	000	80.810	
3 6	200	991.159	0	0000	2000	Ba(Zno. 33 N Do. 67) Os	701.607	810.00	
رم م	341.822	027.750	0 (4.004	047.7	Ba(Zn _{0.83} Ta _{0.67})O ₃	328.145	205.476	
<u>్ల</u>	301.306	557.848	20	4.116	7.173	Ca(Ni, 3,Nb, 1,1)0,	169.700	58.411	
35	351.366	595.036	œ	4.205	7.842	Ca(Ni, TB,)O.	228.688	60.698	
22	246.009	676.138	00	4.160	5.671	DP(CodM)	336.883	65.939	
33	339.078	631.629	00	4.290	7.129	DF(Co).33±1 Co).87) (3	395 871	64 481	
9.6	983.886	616 730	œ	4.256	5.682	FD(C00.33 LE0.67) C3	356.0	2000	
9 6	989.698	2010	·	2011	}	Pb(Mg _{0.33} N b _{0.67})O ₃	020.400	00.800	
2	0.40	270 010	٥	1 956	7 276	Pb(Mg0.33 Tao.67) Us	384.440	04.900	
200	342.049	010.947	0 0	4.200	0000	$Pb(Mn_{0.33}Nb_{0.67})O_3$	335.565	1	
67 	294.009	611.960	ж —	4.240	0.330	$Pb(Ni_{0.83}Nb_{0.67})O_3$	336.810	65.208	
							-		

5.903 7.265 6.526 7.884 6.479 7.935 6.568 6.568 7.828 7.828

4.186 4.181 4.168 4.167 4.086 4.086 4.085 4.085 4.085 4.085 4.088

6.992 6.459 6.340 6.309 4.582 6.948

4.006 4.006 3.945 3.947 4.148 3.945 3.945

∞ ∞ ∞ ~ ∞ 4 4

8.574

4.025

6.554 8.017 6.784 6.784 8.108 7.295 6.511 7.953 4.823 6.254 8.481 8.481 8.883 8.883

4.115 4.074 4.072 4.260 4.260 4.242 4.094 4.091 3.880 3.930 4.040 4.040 4.040 4.040 4.040

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Compound	-		-			
	Formula weight	Volume	8	a', A	Density	Сошро
	A(B, 33 B, 67) Os (cont.,)0 ₈ (cont.)				4N ::4/-0
$p_{h}(N; T_{h}, T_{h})$	395.798	64.481	_	4.010	10.189	Da(Eug. LA Do
Pb(Zn, Nb, S)	339.007	65.939	_	4.040	8.534	Da(Eu. Fa.
Sr(Ca. ", Nb. ",)O.	211.092	205.726	က	4.093	5.110	Bo (Fe Mo
Sr(Ca, "Sb, 67)O3	230.417	545.338	00	4.085	5.611	Be (Fe N)
$Sr(Ca_{s,s}, Ta_{s,s})O_{s}$	270.080	204.170	က	4.083	6.588	D&(E 00.847 D).
Sr(Cd., "Nb. ".)O.	234.957	68.368	_	4.089	5.705	Be/Fee Te.
Sr(Co, 33Nb, 67)O3	217.313	513.922	∞	4.005	5.616	Be (Ad NP.
Sr(Co, 1.,Sb, 17)O3	236.639	510.082	00	3.995	6.161	Ba(Gd - Pa.
Sr(Co, 33 Tao, 47)O3	276.301	190.423	က	3.989	7.226	Ba (Gd. , Be.
Sr(Cu, 33Sb, 67)O3	238.159	503.403	∞ ∞	3.977	6.283	Be (AdSb.
Sr(Fe, 33Nb, 37)O3	216.295	64.192		4.004	5.593	Ba(GdTB.
Sr(Mg, 33Nb, 47)03	205.888	193.651	က	4.011	5.295	Ba (Ho Np
Sr(Mg, 33Sb, 67)O3	225.214	604.358	os.	3.980	5.930	Ha(Ho, 'Pho
Sr(Mgo.33 Teq. 67)O3	264.876	192.301	ಞ	4.002	6.859	Ba(Ho, Ta
$Sr(Mn, 3Nb, 47)O_3$	215.995	i	1	1	1	Ba(In Nb.
Sr(Mn, rTa, rTa)	274.983	1	1	ļ	1	Be (In Da-
Sr(Ni, 1, Nb, 6, 1)O3	217.240	190.081	က	3.987	5.692	Be (In . Pa.
Sr(Ni, 3, Ta, 67) O3	276.228	188.489	က	3.975	7.298	Ba(In. Be.
$Sr(Pb_{0.33}Nb_{0.07})O_3$	266.238	1	١	1	l	Ba(In. Sb.
$Sr(Pb_{\rho_{33}}Ta_{0.67})O_{3}$	325.226	ı	1	1	1 5	Ba(In. Ta.
$Sr(Zn_{0.33}Nb_{0.67})O_3$	219.437	192.818	က	4.006	5.667	Ba(In) II.
Sr(Zno. 33 Tao. 67)O3	278.425	193.119	<u>ო</u>	4.008	7.180	Ba(La, Nb
	$A(B_0^3 +$	$A(B_{0.3}^{3+}B_{0.3}^{5+})O_{3}$				Ba(Lao., Pa
	10000	201010	0	4 215	6 948	Ba(La _{0.8} Re
Ba(Bio. 5N bo. 5) O3	336.281	042.730	0 0	4.010 4.004	8.030	Ba(La _{0.6} Ta
Ba(Bi _{0.5} Ta _{0.5})U ₃	300.302	70.302	· -	4.293	6.333	Ba(Lu _{0.8} Nb
Ba(Ceo.g.N bo.g)O3	370.808	681 479	1 00	4.400	7.228	Ba(Lu _{0.8} Pa,
Ba(Ceo.gFao.g)Og	961.958	66.923	, ,-	4.060	6.480	Ba(Luo.,Ta
Da(Co _{0.8} 1V D _{0.8})O ₃	307 905	528.690	00	4.043	7.734	Ba(Mno.sN)
Da(CC) \$100.5/C3	306.436	1	1	i		Ba(Mn _{0.8} Re
$\mathbf{B}_{\mathbf{c}}(C_{\mathbf{r}} \cdot \mathbf{B}_{\mathbf{c}}, \mathbf{s}) \mathbf{C}_{\mathbf{s}}$	304.436		İ	1	1	Ba(Mn _{0.6} T8
Ba (Cr11: -10:	330,351	1	1	1	1	Ba(Na _{0.8} N)
Ba(Cir. W)O.	309.033	534.633	00	4.058	7.676	Da(Na _{0.8} F ^a
Ba(Dw. Nb. 10.	313.041	600.571	œ	4.218	6.922	Ba(Nd _{0.8} Fe
Ba(Dv Pa)0.	382.088	667.628	00	4.370	1.600	Da(Na _{0.8} 1
Ba (Dy. Th. 1)0.	357,062	623.930	∞	4.272	009.7	Ba(N ₁₀ , sin D
De(D) 0.5 1 10.3 () () () () ()	315.421	598.438	<u>∞</u>	4.213	7.000	Ba(Fro.sing
Ba(Fr. Pa. 10.	384.468	662.143	∞	4.358	7.711	D8(FT _{0.8} F8 D-/D- M-
Ba(Er, 'Be, 1)0,	362.068	583.020	∞	4.177	8.247	D8(Fr _{0.8} 18 D2(D1 N
Ba(Er., Te., 103	359.442	597.586	တ	4.211	7.988	Da(1940.514 Ba/Bh II
Ba(Er, 5U, 5)0	387.983	651.714	<u>∞</u>	4.335	7.906	D9'0rraT)BCT

TABLE 9.2 (cont.)

Compound	Formula weight	Volume	- 2	a', A	Density
	A(B _{0.8} +B _{0.5})O ₈	O ₈ (cont.)			
Ba(En., Nb., 10.	307.771	615.643	∞	4.253	6.639
Ba(En. Pe. 1)0	376.818	677.530	00	4.391	7.386
Ba(Eu., Ta.,)0,	351.792	615.426	œ	4.253	7.691
Ba(Fe, Mo, 10.	261.232	527.514	00	4.040	6.578
Ba(Fe, Nb, 1)0.	259.716	66.923	_	4.060	6.442
Ba(Fe, Be, 1)0.	306.362	521.660	00	4.025	7.799
Ba(Fe. Te. 10.	303.736	66.726	_	4.056	7.556
Ba(Gd, 'Nb, ')O.	310.416	613.258	00	4.248	6.722
Ba(Gd., Pa., 10.	379.463	675.449	œ	4.387	7.461
Ba(Gd, Be, 1)0.	357.083	699.290	00	4.216	7.912
Ba(Gd, Sb, 10.	324.838	601.212	00	4.220	7.175
Ba(Gd, TB, 1)0,	354.437	613.184	œ	4.248	7.676
Ba(Ho, 'Nb, ')O.	314.256	599.930	œ	4.217	6.956
Ba(Ho, Pho.)0.	383.303	605.339	∞	4.306	7.651
Ba(Ho, The 1)0,	358.277	601.639	00	4.221	1.908
Ba(In. Nb. 10.	231.791	567.458	00	4.139	5.425
Ba(In. Oa)O.	280.438	556.223	œ	4.112	9.696
Ba(In, Pa, 1)0,	300.838	635.169	00	4.298	6.290
Ba(In. Re. 1)0.	278.438	563.151	00	4.129	6.566
Ba(In, Sb., 10,	246.213	565.404	00	4.134	5.783
Ba(In, Ta, 1)0,	275.812	567.664	∞	4.140	6.452
Ba(In, U, 1)0,	304.353	618.470	00	4.260	6.535
Ba(La, Nb, 1)0,	301.246	643.759	œ	4.317	6.214
Ba(La, Pa, s)Os	370.293	701.411	<u>∞</u>	4.442	7.011
Ba(La, Re, s)O3	347.893	631.629	00	4.290	7.314
Ba(La, Ta, k)O3	345.267	651.958	∞	4.336	7.033
Ba(Lu, Nb, s)O	319.276	685.116	∞	4.182	7.240
Ba(Lu, Pa, 1)03	388.323	650.813	00	4.333	7.924
Ba(Lug., Tag., 5)03	363.297	586.797	.	4.186	8.222
	259.260	68.067	— (4.083	6.323
Ba(Mn, Re, s)O	305.907	547.343	00	4.080	7.422
Ba(Mn _{0.8} Ta _{0.8})O ₃	255.283	67.718	-	4.076	0.200
$Ba(Nd_0bNb_0b)O_s$	303.911	622.836	x 0 0	012.4	0.400
$Ba(Nd_{0.5}Pa_{0.5})O_{3}$	372.958	690.807	0	024.4	7 7 7 7
Ba(Nd _{0.5} Re _{0.5})O ₃	350.558	616.295	x 0 (4.200	400.
Ba(Nd _{0.8} Ta _{0.8})O ₃	347.932	626.343	×0	4.278	
0	261.146	68.921	_	4.100	6.290
Ba(Pr., Nb., 103	302.245	77.854	_	4.270	0.444
0	371.292	5.9	<u>∞</u>	4.431	7.085
Ba(Pr., Ta., O.	346.266	οó	_	4.270	
	283.244	545.338	20	4.085	0.830
Ba(Rh, U, 10.	355.806		<u> </u>	ا -	۱ _

OTHER PROPERTIES

TABLE 9.2 (cont.)

Density

a', A

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Compound	Formula weight	Volume	13	a', À	Density		Compound	Formula weight	Volume	Z
	A(B _{0.5} B _{0.5})	B3+B5+)O3 (cont.)						A(B _{0.5} B _{0.5})O ₃ (cont.	Os (cont.)	
Ba(Sc., Nb., 6)O3	254.269	69.986	1	4.121	6.031	.~	$Ca(Gd_0bTa_0b)O_3$	257.177	65.541	_
13a(Sco., OSo., 6)O3	302.916	541.742	20	4.076	7.426		$Ca(Ho_0.sNb_0.s)O_s$	216.996	64.912	_
Ba(Sc, 5Pa, 5)O3	323.316	624.807	∞	4.274	6.872	*=	Ca(Hoo.sTao.s)O3	261.017	65.237	٠,
Bu(Sco. , Reo. , O3	300.916	543.938	∞	4.081	7.347	-	Ca(Ino. & No. s) Os	134.631	62.222	٦,
Ba(Sco. Sbo. s)Os	268.691	550.763	00 (4.098	6.479		Ca(Ino. FIRO. S) Ca	909 008	02.301	٦,
Ba(Sco. s.Tao. s)O3	298.290	555.818	x	4.111	7.127		Ca(Lao, 51 00.5) Os	948.000	67.979	٠,-
Ba(Se _{0.5} U _{0.5})U ₃	326.831	611.900	×0 0	4.240	6.092 6.596		Ce (Mr. Te. 10.	208.021	58.851	٠.
Da(Sm _{0.5} 1\O _{0.5})U ₃ B ₅ (Sm D _{6.7})U ₃	376.013	679 618	0 0	4.308	7.347		Ca(Nd, Nb, 10.	206.651	66.368	. –
Ba(Sm. Ta. 1)	350.987	618.252	0 00	4.259	7.539		Ca(Nd, Ta,)O3	250.672	66.371	7
Ba(Tb, 'Nb, ')O,	311.253	75.633	-	4.229	6.831		Ca(Nio. Wo.s)Os	209.358	230.769	4
Ba(Tb, 'Pa', ')O,	380.300	670.611	00	4.376	7.531		Ca(Pro. NPo. s)Os	204.985	669.99	1
Ba(Tl), Ta, ()O.	377.997	596.948	00	4.210	8.409	~	Ca(Pro. & Tao. 8) Os	249.006	66.701	1
Ba(Tm, Nb, O.	316.258	694.399	00	4.204	7.066		Ca(Sco., Reo., s)Os	203.656	242.942	4
Ba(Tm, Pa, 1)0;		656.688	00	4.346	7.792		Ca (Smo. , Nbo. , O3	209.706	65.866	-
$Ba(Tm_{h_1}, Ta_{h_2}, 0)$	360.279	693.976	œ	4.203	8.055		$Ca(Sm_05Ta_05)O_8$	253.727	66.205	_
$Ba(Y_n, Nb_n, 0)$	276.244	74.088	_	4.200	6.189		$Ca(Tb_0.sNb_0.s)O_3$	213.993	65.384	-
Ba(YorPang)O	345.291	662.299	∞	4.359	6.920		$Ca(Tb_0, _8Ta_0, _8)O_3$	258.014	65.383	1
Ba(Y, Re,)O,	322.891	586.797	∞	4.186	7.307		$Ca(Y_0, _{\mathfrak{s}}Nb_0, _{\mathfrak{s}})O_{\mathfrak{s}}$	178.984	65.232	_
Ba(Y, Ta,)O,	320.265	599.717	œ	4.216	7.092		$Ca(Y_0, Ta_0, s)O_3$	223.005	65.232	<u>ر</u>
$B_a(Y_0, U_0, 0)O_3$	348.806	656.235	00	4.346	7.059		Ca(Yb, , Nb, , ,)O3	221.051	64.281	-
$Ba(Yb_0,sNb_0,s)O_3$	318.311	587.217	00	4.187	7.199		Ca(Ybo.sTao s)Os	265.072	64.279	-
Ba(Ybo., Pao., 5)O3	387.358	653.520	00	4.339	7.871		Pb(Fe. &N bo. g)O	329.565	64.819	-
$Ba(Yb_05Ta_05)O_3$	362.332	590.590	00	4.195	8.147	-	Pb(Fe., gTao, s)Os	373.586	64.529	٠,
Ca(Alo. 3Nho. 5)O3	148.022	55.161	H	3.807	4.455		Pb(In ₀ , N _{D₀, s)O_s}	301.641	69.427	<u> </u>
$Ca(Al_{0.5}Ta_{0.5})O_3$	192.043	55.161	_	3.807	5.779	=	Pb(Ho, sNb, s)Os	384.106	71.057	٦,
Ca(Co., 5 W., 6)O3	209.410	235.054	4	3.888	5.917	- ·	FD(Lug. PDg. s) Cg	389.120	70.040	1 -
$Ca(Cr_0{\delta}Mo_{0.\delta})O_3$	162.046	226.583	4	3.841	4.749		Fb(Lu., Ia., s)Os	433.147	00.00	1 -
$Ca(Cr_{0.5}Nb_{0.5})O_3$	160.529	57.061	_	3.850	4.670		FD(Sco. & N Do. s) Os	324.119	07.301	7 -
$Ca(Cr_0. sOs_0. s)O_3$		225.423	4	3.834	6.161		FD(3C0.8.140.8) C3	368.140	71 473	4
$\widetilde{\operatorname{Ca}}(\operatorname{Cr}_0, \mathfrak{s}\operatorname{Re}_0, \mathfrak{s})\operatorname{O}_3$	207.176	225.717	4 -	3.830	6.095		F D(1 D0. 51 D0. 5) O3 Db(Vb	439.182	70.445	-
Ca(Cro.s.Tao.s.)O3	204.550	200.100		00000	6.070		Sr(Co ND 1)	211.538	60.698	
Ca(Cro.s Wo.s)O3	200.001	65 309	* -	0.000	6.0.0	*	Sr(Co, 'Sb, ')O.	225.960	489.304	<u>∞</u>
Ca(L)yo.sIN bo.s)O3	950.001	85 303		070.#	6.45		$Sr(Cr_{s,s}Mo_{s,s})O_{s}$	209.586	478.212	00
Ca(Dyp.s 1 Ep. s) O3	208.802	64.756	٠-	4.016	5.592		Sr(Cr., Nb., 5)03	208.069	61.261	_
Ca(Fr. Te.)O.	262.182	64.757	,	4.016	6.721	-	$Sr(Cr_0, Os_0, \epsilon)O_3$	256.716	481.890	00
Ca(Fe, Mo, 1)O,		231.688	4	3.869	4.699	-	$Sr(Cr_0, Re_0, s)O_s$	254.716	478.212	00
Ca(Fe, Nb, 1)O.	162.455	58.703	_	3.886	4.594	_	$Sr(Cr_0, sSb_0, s)O_s$	222.491	485.958	00
Ca(Fe, 55b, 5)03	176.877	234.551	4	3.885	5.007		$Sr(Cr_0, Ta_0, s)O_3$	252.090	61.163	_
Ca(Fe, ,Ta, ,)O3	206.476	58.701	_	3.886	5.839		Sr(Cr _{0.5} W _{0.5})O ₃	253.541	478.212	<u>~</u>
Ca(Gdo. 8Nbo. 5)O3	213.156	65.540	-	4.032	6.399		$\mathrm{Sr}(\mathrm{Dy_{0.5}Ta_{0.5}})\mathrm{O_{3}}$	307.342	1	
										,

6.514 6.642 3.589 4.751 6.026 6.111 6.111 6.181 6.024 6.197 6.197 6.197 6.197 6.197 6.197 7.924 9.015 10.151 7.924 6.183 6.184

4.032 3.963 3.963 3.963 3.963 3.963 3.963 3.931 4.049 4.049 4.045 4.029 4.029 4.029 4.029 4.029 4.029 4.029 4.029 4.029 4.029 4.029 4.029 4.029 4.039

Density

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Volume

PREPARATION OF PEROVSKITE-TYPE COMPOUNDS

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TABLE 9.2 (cont.)						TABLE 9.2 (cont.)		
Compound	Formula weight	Volume	Z	a', A	Density	Compound	Formula weight	
	A (138 ' B5+10_ (cont.	O. (cont.)					A(B _{0.5} B _{0.5})	
O. T. W. V.	309 722	- (image) #2	ī	-	١	Ba(Cdo., Uo., O.	360.553	
$Sr(Er_0, s.t.w_0, s)O_3$ $Sr(Er_0, s.t.w_0, s)O_3$	302.072	1	1	١	ı	Ba(Coo. & Moo. s)Os	262.775	
Sr(Fe, Mo, 1)03	211.512	491.169	œ	3.946	5.719	Ba(Co., Ke, s)O3	333 890	
$Sr(Fe_{0,k}Nb_{0,k})O_{3}$	209.995	62.571	, m	3.970	5.571	Ba(Co., Co., Co.) C.	306.730	
Sr(Fee, Sbo, k)O3	224.417	496.041	00	3.958	6.008	D8(Co., Wo.,) C3	330.351	
Sr(Feq. 5 Tag. 3) O3	254.016	62.428		3.967	6.754	Ba(Cro. 5 Co. 5) Cr	336.123	
Sr(Ga0., Nb0., 1)O3	216.931	61.443	-	3.946	5.86I	Ba(Fe Re)O.	306.362	
$Sr(Ga_05OB_05)O_3$	265.578	478.212	x 0 0	3.910	1.0.0	Ba(Fe, .U)O.	332.277	
Sr(Ga _{0.6} Re _{0.8})O ₃	263.578	482.444	0 0	3 039	6.319	Ba(Fe, W. 5)O3	305.187	
Sr(Ga., Sb., s)O3	231.333	400.130	-			$Ba(Mg_{\rho,s}O_{g_{\rho,s}})O_{s}$	292.594	
Sr(Gdo. Tao. s)O3	304.717				1	Ba(Mg, Re, 6)O3	290.594	
Sr(Ho, (Ta, s)O,	308.50	66 770	-	4.057	4.527	Ba(Mg., Te, s)O3	261.294	
Sr(Ino. 6N Do. 8) O3	107.011	K92 K07	+ 00	4.030	5.852	Ba(Mg0.5U0.6)O3	316.509	
Sr(In _{0.8} Os _{0.8})O ₃	900.710	59K 7E3	0	4 035	5.777	Ba(Mg0.5W0.5)O3	289.419	
Sr(Ino. & Hou. 6) U.	228.110	700.000	ο α	4 165	6.850	Ba(Mn, 6Reg. 8)O3	305.907	
Sr(In _{0.8} U _{0.8})O ₃	204.033	2003) O	135	6.939	Ba(Mna, U., s)O3	331.822	
Sr(La _{0.8} Ta _{0.8})O ₃	290.047	600.000	5	7.F	3 1	Ba(Ni, Mo,)O	262.663	
Sr(Lu _{0.8} Ta _{0.8})O3	313.577	1 00	0	000	5,516	Ba(Ni, Re, s)O3	307.793	
Sr(Mno. 5 Moo. 5) U3	211.057	000.1.00	0	0.00	6	$B_a(N_i, U_a, 0)$	333.708	
Sr(Mno.,Sbo.,s)O3	223.962	1	Ī	}		Ba(Ni, W, s)O	306.618	
Sr(Ndo. Lao. 6) Os	298.212		1	ł	i I	Ba(Pb, Mo, 1)	336.903	
Sr(Nio.,Sbo.,s)O3	225.848	1 2	<u> </u>	100	6.434	Ba(Sr. Os. 1)0	324.248	
$\mathrm{Sr}(\mathrm{Rh}_{\mathfrak{d},\mathfrak{s}}\mathrm{Sb}_{\mathfrak{d},\mathfrak{s}})\mathrm{O}_3$	247.946	255.808	# 0	0.000		Ba(Sr. Re. 1)O3	322.248	
Sr(Sco. 6 Oso. 6) O3	253.190	010.000	0 0	4.010	6.467	$Ba(Sr_{05}U_{05})O_3$	348.163	
Sr(Sco. 1100. 6) O3	201.190	010.010	9		; ;	Ba(Sro. 8 Wo. 8)O3	321.073	
Sr(Sm _{0.6} Tu _{0.6})O ₃	910 650		1	ł	}	$Ba(Zn_05Os_05)O_3$	313.123	
Sr(1m _{0.5} 18 _{0.8})O ₃	919 819		1	1	1	$Ba(Zn_05Re_05)O_3$	311.123	
Sr(X Do. 6 T Bo 8) Un	910.916	_	_		_	Ba(Zno., Uo., O	337.038	
	A(B2	$A(B_{0.5}^{1+}B_{0.5}^{1+})O_3$				Ba(Zno., Wo., s)Os	309.948	
B./B. O. 10	349.108	625.463	∞	4.276	7.412	Ca(Cao.sOso.s)Os	203.218	
Da(Da ₀ , \$0.50, \$)	347.108	623.271	00	4.271	7.396	Ca(Ca _{0.5} Re _{0.5})O ₃	201.218	
D&(D&0.61.000.5) 03	373 023	702.595	00	4.445	7.051	Ca(Ca _{0.8} W _{0.8})O ₃	200.043	
D2(D20.500.5) 03	345 933	636.056	00	4.300	7.223	Ca(Cd0.5Re0.5)O3	237.378	
D&(D&0.5 W 0.5) O3	953 348	683.229	00	4.177	5.769	$Ca(Co_0.sOs_0.s)O_3$	212.645	
DE (CEO. SINTO S. S.)	200.00	584.696	00	4.181	6.825	Ca(Coo., Reo., O	210.645	
Da(Ca ₀ , sOB ₀ , s)O ₃	900.478	583 439	00	4.178	6.194	$Ca(Fe_{0.5}Re_{0.5})O_{8}$	209.102	
Ba(Ca _{0.5} IXe _{0.5})O ₃	960.170	501 993	· 00	4.196	6.046	Ca(Mg _{0.8} Re _{0.8})O ₃	193.334	
13a(Cao. 5 Teo. 5)U3	204 203	651 714) oc	4.335	0.610	Ca(Mg0.6W0.5)O3	192.159	
Ba(Ca _{0.5} U _{0.6})U ₃	324.333	590 590	000	4.195	6.685	Ca(Mn _{0.8} Re _{0.8})O ₃	208.647	
Ba(Ca _{0.8} W _{0.8})O ₃	297.303	576 969) oc	4.162	7.748	Ca(Ni _{0.8} Re _{0.8})O ₃	210.533	
Ea(Cd _{0.8} Os _{0.8})O ₃	334 638	576.346		4.161	7.711	Ca(Sr _{0.8} W,s)O ₈	223.813	
D&(Cu _{0.6} 1ve _{0.6}) C ₃			_					

7.734 7.649 7.649 7.646 7.681 7.684 7.684 7.684 7.684 7.369 7.369 7.236

4.316 4.043 4.043 4.043 4.049 4.040 4.040 4.060 4.060 4.060 4.060 4.060 4.060 4.060 4.060 4.060 4.060 4.060 4.060

∞ ∞ ∞ ∞ ∞ ∞ ∞ ∞ ∞ ∞ ∞ ∞ ∞ ∞ ∞ ~ ∞ ∞ ∞

537.963 527.514 527.906 537.368 588.691

571.167 591.506 521.660 574.271

321.487 66.081 528.690 587.217 531.047

B2+B6+)O3 (cont.,

647.343 618.470 65.086

531.244

579.259 524.777

519.718

619.686 613.128 690.807

614.125 530.457

532.623

6.949 6.980 6.693 6.943 7.7839 7.7839 7.760 7.760 7.760 6.062 6.062 6.035 6.03

4.263 4.248 4.248 4.250 4.047 4.063 4.040 4.060 4.000 4.000 3.897 3.860 3.897 4.050

263.819 512.000 260.017

235.445 234.899 230.064 236.743

456.533

592.069 534.596 261.552

156 PREPARATION OF PEROVSKITE-TYPE COMPOUNDS

ont.)
9.2 (
TABLE

A(B _{0.5} B _{0.5})O ₃ (cont., 367.153 —
367.153
403.313
331.144
359.269
203.628
250.758
248.758
274.673
247.583
210 600
913 OKK
260.185
258.185
284.100
257.010
280.631
269.313
258.642
256.642
282.557
100.H01
949.874
240.874
211.574
266.789
239.699
256.187
282.102
255.012
212.943
258.073
283.988
256.898
287.183
274.528
272.528
298.443
)
271.353

TABLE 9.2 (cont.)

Compound	Formula weight	Volume	Z	a', À	Density
!	A(B\$+B6+	A(B _{0.5} ² +B _{0.5} ⁰ +)O ₃ (cont.)			
$Sr(Zn_{\mathfrak{d}.\mathfrak{s}}W_{\mathfrak{d}.\mathfrak{s}})O_{\mathfrak{s}}$	260.228	502.438	<u>∞</u>	3.976	828.9
; ;	A(B _{0.5}	A(B;;B;;)O3			
Ba(Ago., I.o., s)O.	302.725	605.496	∞	4.230	6.640
Ba(L10.8080.8)O3	283.908	531.441	∞	4.050	7.094
Ba(Lio. Reo. 5)03	281.908	534.992	∞	4.059	6.998
Ba(Na _{0.5} I _{0.5})O ₃	260.285	578.010	ò	4.165	5.980
$Ba(Na_0.sOs_0.s)O_s$	291.933	567.869	∞	4.140	6.827
$\mathrm{Ba}(\mathrm{Na_{0.8}Re_{0.8}})\mathrm{O_3}$	289.933	570.961	00	4.148	6.744
$Ca(Li_0, Os_0, \epsilon)O_s$	186.648	480.049	00	3.915	5.163
Ca(Lio. Req. 6)03	184.648	480.049	00	3.915	5.108
Sr(Li _{0.5} Os _{0.5})O ₃	234.188	485.588	00	3.930	6.405
Sr(Lio.5Reo.5)O3	232.188	487.443	œ	3.935	6.326
Sr(Na _{0.5} Os _{0.6})O ₃	242.213	537.368	00	4.065	5.986
$\mathrm{Sr}(\mathrm{Na_{0.8}Re_{0.8}})\mathrm{O_3}$	240.213	537.368	∞	4.065	5.936
	A*+(B2+	+B(+)03			
La(Co, Lr,)O,	216.375	. 1	1	i	
$\mathrm{La}(\mathrm{Cu}_{0.8}\mathrm{Lr}_{0.8})\mathrm{O}_3$	218.678			ļ	1
La(Mgo., Geo., s)O3	235.359	59.319	7	3.900	6 586
$\mathrm{La}(\mathrm{Mg_{0.5}\mathrm{Ir_{0.6}}})\mathrm{O_3}$	199.064	496.793	∞	3.960	5.321
La(Mgo.sNbo.s)Os	245.517	1	I	ł	
La(Mgo., Ruo., 6)O3	249.699	494.914	∞	3.955	6.697
La(Mgo.sTio.s)Os	223.014	60.791	-	3.932	6.090
La(Mn _{0.8} Ir _{0.8})O ₃	214.377	485.588	œ	3.930	5.863
La(Mno. & Ruo. s) Os	264.912	481.890	œ	3.920	7.300
La(Nio. 5Iro. 5)Og	216.263	493.039	<u>∞</u>	3.950	5.825
La(Nio. 8 Ruo. 8)O3	266.798	493.039	00	3.950	7.186
La(N _{10.5} 11 _{0.8})O ₃	240.213	60.698		3.930	6.569
La(2n _{0.5} Fu _{0.5})O ₃	270.128	292.909	x -	3.985	7.086
114(14E0.8140.5) 08		810°60	-	0.800	0.030
	₩.	B.,7,3)O3			
Ba(Na _{0.25} Ta _{0.75})O ₈	326.797	70.804	_	4.137	7.662
Sr(Na _{0.25} Ta _{0.75})O ₃	277.077	66.676		4.055	6.898
	A(B3+B	A(B _{0.5} B _{0.5})O _{2.75}			
$\mathrm{Ba}(\mathrm{In}_{0.6}\mathrm{U}_{0.6})\mathrm{O}_{3.76}$	300,353	625.246	<u>∞</u>	4.275	6.379
	$A(B_{0.5}^{2+}E$	$A(B_{0.5}^{2}+B_{0.5}^{6}+)O_{2.75}$			
Ba(Ba, 18a, 5)O3.75	340.482	656.235	<u>∞</u>	4.345	6.890
Ba(Feo. 8 Mou. 8) O3. 75	257.232	527.514	00	4.040	6.476
C (E ")"	985 009	700 084	a	7	1000

There also is a small amount of mechanical property data available on perovskite solid solutions of the PZT type, Pb($Zr_{1-x}Ti_x$)O₃.⁽⁵⁾ The interest in these materials as piezo-electrics was discussed in Chapter 5. Tensile strength measurements on hot-pressed specimens of Pb($Zr_{0.45}Ti_{0.48}$)O₃, Pb($Zr_{0.45}Ti_{0.45}$)O₃ and Pb($Zr_{0.45}Ti_{0.48}$)O₃, Pb($Zr_{0.45}Ti_{0.45}$)O₃ and Pb($Zr_{0.45}Ti_{0.48}$)O₃ containing 1 wt%, Nb₂O₅ balanced with PbO gave values of 9700, 11,150, 12,550 and 14,600 psi respectively. The nominal grain size of these particles was 3 microns and the densities of the specimens were at least 95% theoretical. The moduli of the hot-pressed specimens of Pb($Zr_{0.45}Ti_{0.45}$)O₃ and Pb($Zr_{0.45}Ti_{0.45}$)O₃ described above were found to be 14×10° psi for the former two phases and 16×10° for the latter phase.

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CHAPTER 10

PREPARATION OF PEROVSKITE-TYPE OXIDES

10.1. POWDERS

nium silicate boats or in platinum crucibles. However, the material often will react with the platinum or impurities in addition, since the oxides often can not be entirely reacted mixed thoroughly and fired at 1000°C for 24 hr in zircofore, if extremely pure material is desired it is advantageous to fire the mixture on top of a barium titanate compact. In by a single firing, repeated regrinding and reheating is re-Many of the perovskite-type oxides can be prepared by a high-temperature solid-state reaction between binary oxide ides, if they can be obtained in smaller particle sizes to insure powders which are stable in air. However, it is often advantageous to use carbonates, nitrates, etc., instead of the oxquicker reactions or if they are much purer. In a typical preparation of barium titanate powder, barium carbonate and titanium oxide are weighed out in equal molar quantities, the boats as evidenced by coloration of the sample. There-

Some of the perovskite-type compounds can only be prepared using special techniques. These are discussed below.

Compounds containing lead in the A position of the perovskite structure are often difficult to prepare because of the volatility of lead oxide. Sometimes this problem can be alleviated by heating the reactants in a lead oxide atmosphere, by using an excess of lead oxide in the reaction mixture, or by heating the reactants at a relatively low temperature to allow them to combine before the final firing. When lead is the A ion and pentavalent ions are used as the B ions,

compounds with the pyrochlore structure very often are formed early in the reaction sequence and are difficult to react further even with repeated mixing and reheating of the

Fresia et al. (1) found that in the preparation of compounds with the general formula $A(B_0^2;W_{0,5})O_3$, where A=Sr, Ba, $B^{2+}=Fe$, Co, Ni and Zn, alkaline earth tungstates always could be detected in the final products. Similar results have been observed for molybdenum compounds. Regrinding and refiring of the samples often helps to reduce the amount of tungstate or molybdate present, but each compound has to tungstate or molybdate present, but each compound has to atture and firing time must be determined.

In preparing perovskite-type compounds containing divalent iron and cobalt, the valence state is retained by heatling the sample in an evacuated sealed silica capsule or in a non-oxidizing atmosphere. Divalent Fe or Cr ions can be non-oxidizing atmosphere. Divalent Fe or Cr ions can be obtained in a compound by mixing equal amounts of metallic Fe and Fe₂O₃ or Cr and Cr₂O₃, respectively, with the other oxide constituents. For example, Galasso et al. (2) prepared oxide constituents according to the equation:

 $B_{aO} + \frac{1}{9}Fe_{2}O_{3} + \frac{1}{9}Fe + \frac{1}{3}Ta_{2}O_{5} ---- Ba(Fe_{0.33}Ta_{0.67})O_{3},$

compacting the sample and firing it at 1000°C for 24 hr

in an evacuated sealed silica capsule. Sleightand Ward⁽³⁾ informing compounds of the $A(B_0^2 + U_0^4 + J)O_3$ Sleightand Ward⁽³⁾ in forming compounds of $B_2 + B_2 O_3$ as a source type found it advantageous to use $UO_2(NO_3)_2 \cdot 6H_2 O$ as a source of hexavalent uranium. Compounds containing pentavalent uranium were prepared by heating UO_2 and UO_3 in equal proportions with the other oxide reactants. The UO_3 was proportions with the other oxide reactants. The UO_3 was obtained by heating $UO_2(NO_3)_2 \cdot 6H_2 O$ at $400^{\circ} C$ in air and the UO_2 was prepared by heating UO_3 at $1000^{\circ} C$ in hydrogen.

Sleight et al. (4) also produced other complex perovskite compounds such as Sr(Na_{0.5}Re_{0.5})O₃ and Sr(Na_{0.5}Os_{0.5})O₃ which contained heptavalent rhenium and osmium by heating the metal with sodium carbonate and strontium oxide in air. A similar procedure was used to prepare compounds with hexavalent osmium as one of the B ions. The reaction

is given as:

$$AO + \frac{1}{2}BO + \frac{1}{2}Os + \frac{3}{4}O_2 \longrightarrow A(B_0.\epsilon Os_0.\epsilon)O_3$$

where A = Sr or Ca, and B = a divalent metal ion.

A mixture of (ReO₃+Re) was used in the preparation of compounds containing pentavalent rhenium with other trivalent ions in the B position.

Patterson et al. (5) found that they could prepare A(B3; W5; V0; and A(B0,5)M0,5)O3 type compounds by using a mixture of the metal trioxides and the metals to obtain pentavalent tungsten and pentavalent molybdenum respectively.

Ridgley and Ward⁽⁶⁾ prepared the strontium-niobium Ridgley and Ward⁽⁶⁾ prepared the strontium-niobium bronzes which contained some niobium ions in the tetravalent state. These phases adopted the perovskite structure when x varied between 0.7 and 0.95 in Sr_xNbO₃. Two procedures were used to prepare these bronzes. The first involved the following reaction:

(0.5+x)SrO+0.4xNb+(0.5-0.2x)Nb₂O₅------Sr(0.5+x)NbO₃

where Nb metal was used to reduce part of the pentavalent, niobium and the second required that NbO₂ be formed first and reacted with SrO and Nb₂O₅. The NbO₂ was prepared by reducing Nb₂O₅ with hydrogen at 1200°C for 36 hr.

Other methods for producing compounds containing elements in unusual oxidization states have been reported by McCarroll et al. ''Randall and Ward' and Kestigian et al. '0' McCarroll et al. prepared CaMoO₃ using a mixture of molybdenum num metal and MoO₃ as a source of tetravalent molybdenum, num metal and Ward prepared SrRuO₃ by heating strontium Randall and Ward prepared SrRuO₃ by heating strontium oxide and ruthenium metal in air, and Kestigian et al., prepared LaVO₃ by mixing and heating La₂O₃ and V₂O₃ together in vacuum. The V₂O₃ was obtained by heating vanadium pentoxide in hydrogen at 800° for 14 hr.

penuoxius in hymres... oxides which are unstable be-Many of the ions in binary oxides which are unstable become stabilized in the perovskite structure. For example, BaFe⁴+O₃ can be prepared although heating Fe₂O₃ in oxygen will not produce Fe⁴⁺. As another example, the addition of a vill not produce Fe⁴⁺. As another example, the addition of a rare earth oxide or Y₂O₃ to barium titanate produces some Ti³+ when the mixture is heated to 1000°C in air. This material is extremely stable, even though heating Ti₂O₃ in air PREPARATION OF PEROVSKITE-TYPE OXIDES

would result in the formation of TiO_2 . This factor is used to advantage in the preparation of many of the perovskite type compounds.

The preparation of the magnetic perovskites BiMnO₃ and BiCrO₃ created an interesting new area of research on perovskite type compounds. ⁽¹⁰⁾ These compounds were formed by heating the oxides at 700°C under a pressure of 40 kbar, and then quenching them. In both cases, they formed compounds which had distorted perovskite structures with triclinic unit cells.

10.2. THIN FILMS

The need for special elements in microcircuitry has caused considerable interest in the formation of thin films of dielectric materials. The perovskitc-type compounds which have high dielectric constants are most attractive for this purpose. However, the problems of forming binary oxides in thin film form have not been entirely solved, thus, researchers have been reluctant to attempt the preparation of thin films of more complicated materials. Of the studies which have been conducted on perovskites, most of them have been on BaTiO₃ films. Films 7.5 μ thick have been formed by a special slip method⁽¹¹⁾ but thinner films could not be prepared. In addition, thin single crystal films have been prepared by first growing crystals from solution⁽¹²⁾ and then etching in hot phosphoric acid.⁽¹³⁾ Films also have been obtained by spreading small amounts of molten BaTiO₃ on a platinum sheet.⁽¹⁵⁾

approximately 1 to 2 μ thick by vapor deposition. (49) In this approximately 1 to 2 μ thick by vapor deposition. (49) In this process, the BaTiO₂ powder was mixed with alcohol, placed on a tungsten-wire filament and vacuum evaporated onto a platinum substrate. The BaO evaporated first and the TiO₂ plater, but after firing the film at 1000–1100°C the barium titalater, but after firing the film at 1000–1100°C the barium titalater, but after firing a gain formed. X-ray diffraction studies indicated that the films consisted of mainly BaTiO₃ with traces of BaO₂ and TiO₂ and with minor amounts of BaTi₄O₉, BaTi₃O₇, and Ba₂TiO₄. The samples were prepared for property measurements by placing a gold dot on the surface to act as an elec-

trode. The films were found to be ferroelectric and had dielectric constants as high as 270. This, of course, is one of the simpler techniques of forming barium titanate films, but is not satisfactory for obtaining pure materials.

Using a more elaborate method, Green⁽¹⁷⁾ deposited alternating layers of BaO and TiO₂ by successive evaporation from several tungsten coils and then heating the films in air at 1150°C. Frankl et al.⁽¹⁸⁾ used two electron beams to evaporate TiO₂ and BaO and Moll⁽¹⁹⁾ evaporated barium titanate in an electric field to obtain single-crystal titanate films; however, Roder's studies⁽²⁰⁾ left some doubt as to the reliability

before evaporation resulted in films with lower dielectric constants, 250-300 for Ba(Tio, Sno,1)O3 and 200-400 for $\mathrm{Sr}_{0.73}\mathrm{Ba}_{0.27}\mathrm{TiO}_3$. The dissipation factor at 1 kc/s for $0.2~\mu$ films was larger for crystalline films of barium titanate than for the amorphous films and the breakdown strength of the amorphous films exceeded 1.5×10 V/cm while that of the crystalline films was 2 to 3 times lower. However, the dielecbut using a substrate temperature of 500°C resulted in the Substitution of strontium or tin into the barium titanate tric constant of the amorphous films was only from 13-20. ted through a rod (see Fig. 10.1). The grains were dropped onto tained and from which the material was evaporated onto a ture was held 2300°C for the BaTiO₃ evaporation, slightly lower for those containing some BaSnO₃. When the substrate formation of crystalline films. The crystalline films of BaTiO₃ were found to have a dielectric constant of 400 to 700. an iridium boat where a liquid pool of the material was mainsubstrate held at approximately 500°C. The boat temperaform by the same technique. The samples were formed into pellets crushed and sieved to form grains of 100/200-mesh size. The grains then were delivered by a V-shaped niobium trough and were moved by taps from a gear wheel transmitwas held at room temperature, amorphous films were formed, titanate was used by Müller et al. (21) who prepared thin films grain by grain of powder. In addition, solid solutions of BaTiO3 with SrTiO3 and BaSnO3 were prepared in thin film The best technique for producing thin films of barium with thicknesses of the order of 1 μ by vacuum evaporating of Moll's process.

from the boat before the next one arrives, but the technique of using a molten pool was preferred by Müller because it reduced contamination from the boat and permitted the use It should be noted that the films also were prepared using a flush evaporation technique where the grain is evaporated of lower heating temperatures.

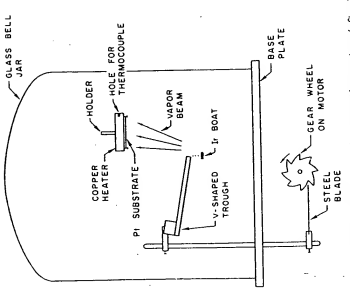


Fig. 10.1. Diagrammatic view of evaporation system (after Mülller constant).

grain evaporation technique. (22) A cleaved LiF substrate was rigidly fastened to a copper block and heated to 700°C. The Müller et al. also deposited thin films of SrTiO₃, CaTiO₃, BaSnO3, SrSnO3, BaCeO3 and NaNbO3 using the grain by results of these experiments indicate that the technique may be feasible for producing single crystals of these materials.

Attempts also have been made to produce thin films of PbTiO₃ with tolerances of $\pm 0.1~\mu$ by generating a plasma of

a dielectric constant of over 100. The breakdown voltage of films on a low temperature substrate. (23) Thin films have been formed with capacitances of the order 50 $\mu \mathrm{F/in^2}$ indicating the bulk material in a vacuum chamber and forming the thin the thinnest films has been found to be 10 volts.

10.3. SINGLE CRYSTALS

structure, those with ferroelectric and magnetic properties and those with potential application as laser host materials Of the large number of compounds with the perovskite have received the most attention from single-crystal researchers.

KNb03

Single crystals of potassium niobate, KNbO₃ and BaTiO₃ have been studied most extensively. Matthias and Remeika (24)

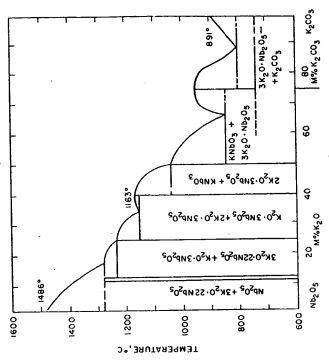


Fig. 10.2. Phase diagram of the $\rm K_2CO_3-Nb_3O_5$ system (after A. Reisman and F. Holtzberg, J. Am. Chem. Soc. 77, 2117 MOL % (1955).

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tallization resulted in clearer and less colored crystals. The problem with the flux technique in general is that the crucible is often attacked and the crucible material and solvent is shown in Fig. 10.2. Pulvari(27) evaluated the methods of these previous workers and tried additional fluxes and flux combinations to obtain pure crystals. The fluxes used were $\rm K_2CO_3,~KCl,~NaCl,~KF, ^{'}CaCl_{2'}$ $^{'}KBO_2,~K_2SiO_3$ with $\rm K_2CO_3$ being selected as the most satisfactory one for growing ferroelectric crystals. The purest crystals were grown with 5-10%excess of $m K_2CO_3$ and a soak time in the $1080-1100^{\circ}C$ temperature range before slowly cooling the melt. Repeated recryswasser(26) also grew $\mathrm{KNbO_3}$ from a flux. The phase diagram have reported the growth of KNbO₃ single crystals using KCl or KF as a flux. In later studies, Shirane et al. (25) and Triebenters in the growth process.

$NaNbO_3$

at 5°/hr. The crystals prepared in this manner grew in the a mixture of sodium carbonate and niobium pentoxide in a sodium fluoride flux. (28) In this process, the mixture is preheated to 1000°C, soaked for 2 hr at 1350°C and then cooled Sodium niobate, NaNbO₃, crystals have been grown from form of small cubes.

NaTaO₃

Na₂B₄O, and Ta₂O₅ in the proportions 7:1:4 is heated at gives the following method for growing single crystals of sodium tantalate, NaTaO3. A mixture of Na2CO3, [200°C for 12 hr and then cooled over a period of 6 hr resulting in crystals whose dimensions are $1\times2\times2$ mm. $\mathrm{Kay}^{(29)}$

KTaO3

water and are then used as seeds to pull clear $\mathrm{KTaO_3}$ crystals from a melt. These blue crystals are not adequate for ferroand then cooled more quickly to room temperature. The dark using a KF flux (30) In a typical run, a mixture with a flux to sample ratio of 5:1 mole % is melted in a platinum crublue-black crystals produced are leached from the flux with Potassium tantalate, KTaO3, crystals have been prepared cible, soaked at $> 1300^{\circ}$ for 4 hr, cooled at 30° /hr to 900° C, electric applications since they are highly conducting

tinum crucible, and set into a vertical furnace. An oxygen Fig. 10.3). The mixture is slowly heated to a temperature a typical charge of 70 g of Ta2Os, 42.10 g of K2COs, 24 mg of MnO₂ and 12 mg SnO₂ are mixed, placed in a 100-ml plaatmosphere is maintained in the furnace (see phase diagram, In the process used by Wemple(30) to pull crystals of KTaOs,

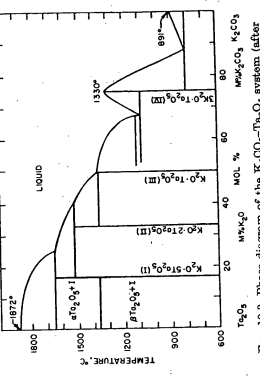


Fig. 10,3. Phase diagram of the K₂CO₃-Ta₂O₅ system (after A. Reisman, F. Holtzberg, M. Berkenblit and M. Berry, J. Am. Chem. Soc. 78, 4514 (1956)).

lifted above the melt surface but still kept in the furnace, and is lifted 1-2 mm. When the crystal is at the desired size, it is 0.5 cm from the melt surface, the temperature of the melt is a temperature 2-3°C above T_L , the seed is lowered to touch the melt and lifted 1-2 mm, pulling a small meniscus. The seed drive motor then is set to rotate at 60 rev/min with reversal every 30 sec. During the growth period the cooling rate is maintained at 3.5°C/hr and at various times the seed the cooling rate is changed to 25-30°C/hr until room temsoaked for 4-6-hr. The KTaO₃ seed then is lowered to within cooled at the rate of 3.5°C/hr. When the melt passed through 10-20°C above the liquidus over a 12-15-hr period and raised 5°C above the liquidus temperature, T_L , and slowly

PREPARATION OF PEROVSKITE-TYPE OXIDES

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perature is attained. The color of the crystals grown in this manner changed from a bright green to colorless at room temperature and the crystals weighed from 4-10 g.

KTN

was selected from the phase diagram so that crystals of electrical losses and a large saturation polarization. A mixture of K₂CO₃, Ta₂O₅, Nb₂O₅ and SnO₂ in appropriate proportions to obtain a composition $K_{1+x}(Ta_{0,29}Nb_{0,71})O_{8}Sn_{0,001}$, where x crucible and set in a vertical tube furnace. The composition exhibit a large room-temperature electro-optic effect, low depends on volatilization losses, were placed in a platinum K(Ta_{0.63}Nb_{0.37})O₃ would be pulled from the melt (see Fig. 10.4). The crucible was placed on a pedestal which was rotat-A similar procedure was used by Wemple 800 and Bonner et al. (31) to grow crystals of KTN, K(Tao. KDo.37)Os, which

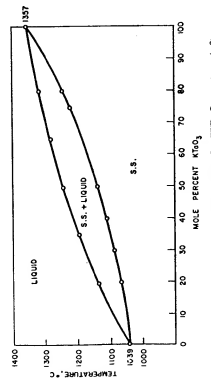


Fig. 10.4. Phase diagram of the KNbO_s-KTaO_s system (after A. Reisman, S. Triebwasser and F. Holtzberg, J. Am. Chem. Soc. 77, 4228 (1955)).

muffle which contained the crucible was closed by a split The furnace was held at approximately 1225°C, oxygen flow was maintained up through the muffle, and the bottom of The melt was then cooled 0.1°C/hr while the seed was rotated at 60 rev/min with reversals every 30 sec. The top of the the crucible was kept 50°C warmer than the top of the melt. ring with a hole to accommodate the rod that held the seed.

at 1/2 in. per day until it was cool enough to remove from the from 2 to 3 days, and then the growing crystal was lifted ed at the surface. The seed was allowed to grow laterally furnace. It is difficult in this process to produce a crystal with a uniform composition throughout.

BaTiO,

tals have been studied extensively. Crystals of BaTiOs Because of its ferroelectric properties, BaTiO_s single cryshave been grown from fluxes, by the Czochralski technique, the Verneuil technique and also using zone melting methods

Some of the fluxes used to grow BaTiO₃ include BaCl₂, ⁽³²⁾ twins (shaped like butterfly wings) and cubes using BaF₂ as KF(38) and BaF, (34) Linares obtained small blue butterfly

these butterfly twin crystals and concluded that they were tals being formed. DeVries (35) conducted detailed studies on probably formed when (111) twinning of the (100) habit takes mentioned fluxes for growing BaTiO, single crystals. In this technique, a mixture of 10 mole % BaTiOs and 90 mole % KF is soaked at temperatures of 980 to 1200°C for 4 hr followed by cooling rates of 10°C/hr, resulting in butterfly orys-Potassium fluoride is a more popular flux than the aboveplace.

the surface with air flowing through the seed rod. After 30 min the melt was cooled 5°C/hr for 1 hr. The crystal then was pulled at 0.25 mm/hr and the melt was cooled at 2 to 3°C/hr. The seed was rotated at 60 rev/min and reversed at 30-sec intervals. Then, the crystal was removed from the melt at 1335°C and annealed to room temperature. Single crystals of BaTiO₃ up to 2 cm diameter by 1 cm long have from a BaÓ-TiO₂ melt by von Hippel et al. The phase diagram is shown in Fig. 10.5. A detailed schematic of the cryswas made up of 67 mole % TiO2 and 33 mole % BaO. The mixture was soaked at 1420°C, the temperature was lowered to 1396° and the seed then was immersed about 1 mm below tal growth furnace is shown in Fig. 10.6. The mixture used Single crystals of BaTiO₃ also have been grown by pulling been grown by this method.

Attempts also were made by von Hippel et al. (38) to grow single crystals of BaTiO₃ by the flame-fusion technique. For this technique powders with small particle size but with good flow characteristics are necessary. In order to produce the powder, a solution of titanium tetrachloride, prepared by dripping 1.50 moles of TiCl₄ into 500 ml of water below 20°C, was added to a solution of oxalic acid, 5 moles (COOH)₂ in 1320 ml of water and held at 20°C. A solution of barium chloride at 70°C, 1.60 moles BaCl₂·2H₂O dissolved in 900 ml

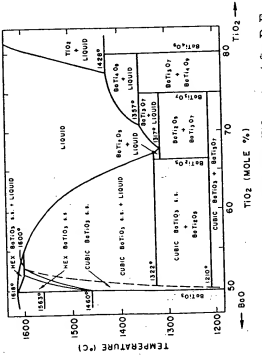
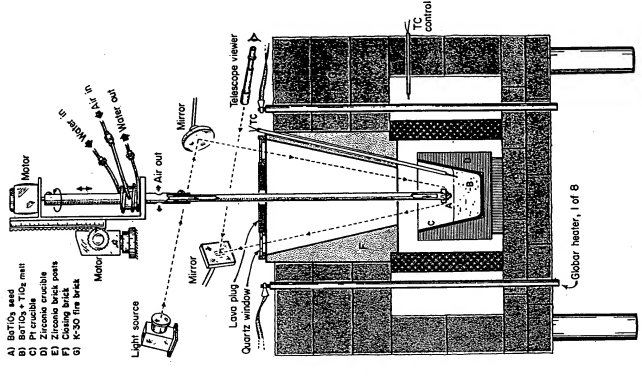


Fig. 10.5. Phase diagram of the BaO-TiO, system (after D. E. Rase and R. Roy, J. Am. Geram. Soc. 38, 110 (1965)).

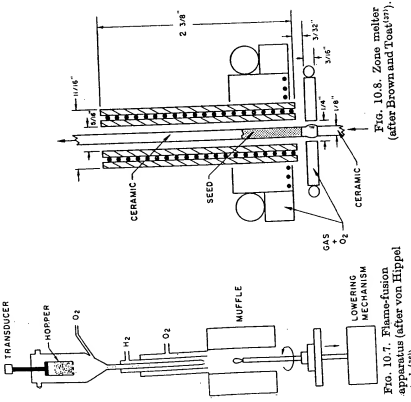
H₂O, was added to the mixture with rapid stirring. After 5 hr of stirring, barium titanyl oxalate was filtered, washed and ignited for 2 hr at 1000°C. Powder was passed through a sieve, and was used in the flame-fusion apparatus (see Fig. 10.7). Small crystals of BaTiO₃ were obtained, but they were not of optical quality.

Brown and Toat⁽³⁷⁾ found that they could grow single crystals 2.5 cm long by 0.32 cm diameter of BaTiO₃ which contain 1.5%, SrTiO₃ by a floating zone process. The heating source was a ring burner closely surrounding the molten zone.



Frg. 10.6. Crystal-pulling furnace (after von Hippel et al. ⁽¹⁸⁶⁾).

Pressed bars of the material were fused to a seed crystal and were withdrawn at a rate of 3.5 cm/hr into an auxiliary heater maintained at temperature of 1300–1500°C (see Fig. 10.8).



et al. (36)).

through a steep gradient. The center of the furnace coil was hr. In addition, he grew large grains of BaTiO₃ by seeding a In an interesting study, DeVries (38) obtained large grains of barium titanate in a polycrystalline rod by lowering it only at 1300° C and the rod was lowered at the rate of $0.6\,\mathrm{mm}/$ polycrystalline aggregate.

Of these methods for growing barium titanate, the KF flux method is probably the most successful, since other tech-

highly strained. The floating-zone process produces crystals both require the use of a crucible which may dissolve and be powders must be prepared and the resulting crystals are with less strain than the Verneuil technique, but with more defects than crystals grown by flux and Czochralski techintroduced into the crystal. In the Verneuil technique special tal through a phase transition which causes the crystal to crack, but there are problems with all of the crystal-growing techniques. The flux technique and the Czochralski method od involve relatively rapid and uneven cooling of the crysniques such as pulling from the melt and the Verneuil meth-

CaTiO3

niques.

solutions were soaked at 1150°C and the carbonates at 1000°C for 30-40 hr. Cooling rates used varied from 50°C to 100°C/hr. The crystals produced were of the order of 1 mm. (39) BaCl2, CaCl2-BaCl2 and Na2CO3-K2CO3 fluxes. The chloride Single crystals of CaTiO₃ have been obtained using CaCl₂,

sieve, the powder was placed in the flame-fusion apparatus ter growth, but this was overcome by annealing and slowly lutions of titanium tetrachloride, calcium chloride and oxalic acid in molar proportions of 1.0/1.4/4.0. Then the calcium titanyl oxalate formed was transformed to CaTiO₃ by heating to 1000°C. After passing the CaTiO₃ through a 100-mesh and the crystal was grown. Initially, the boules fractured afmeter, have been grown by Merker using the flame-fusion technique. (40) The feed material was prepared by mixing so-Larger crystals of CaTiO₃, 25 mm long and 12 mm in diacooling the crystal.

SrTi0,

sieved and used. The crystal as prepared had an opaque black process reported for preparing BaTiO₃ boule powder. A soluappearance; however, a colorless transparent crystal was ob-Merker. (41) The feed material was formed by nearly the same followed by the addition of a solution of strontium chloride. After aging, the crystal salt was filtered, heated at 1000°C, Strontium titanate crystals also have been prepared by tion of oxalic acid was added to dilute titanium tetrachloride, The solution temperature was held at 70°C under agitation. ained by annealing the crystal in air.

PREPARATION OF PEROVSKITE-TYPE COMPOUNDS 174

clear crystals 5×3×0.2 mm of PbTiO₃ using a Bridgman-Stockbarger method with excess PbO. Nomura and Sawada Single-crystal growth of PbTiO₃ is of interest because PbTiO3 is a high-temperature ferroelectric. Rogers(42) grew also obtained similar results using a PbCl₂ flux. (43)

Single crystals of CdTiO₃, 0.12×0.06.0.06 mm, have been grown from a NaCl flux. (44)

crucible, heated it at 1250°C for 1 hr and cooling at a rate of 50°/hr. The crystals grown were in the shape of cubes, reported by Jona et al., (46) which involved placing a mixture of 2.4 g of PbF₂ and 6.9 g of PbZrO₃ in a covered platinum Pulvarius grew crystals of PbZrO₃ using the techniques 3 mm on edge.

(R.E.) BO3

perature was then reduced to 850°C at a rate of 30°/hr and the crystals were leached from the mixture and crucible A constituent oxide to lead monoxide ratio of 1:6 by weight was mixed and a platinum crucible was used to hold the mixture. The mixture was maintained at a temperature of 1300°C for a short period, except for compounds containing Al3+ or Sc3+ where the soaking time used was 4 hr. The tem-Fe, Co and Ga in the B position were grown by Remeika. (47) Pr, Nd, Eu, Sm and Gd in the A position and Al, Sc, Cr, A large number of small perovskite crystals with Y, La, with hot dilute nitric acid.

LaAlO3

bits at 435°C presents a problem for growing it as a single Because of the longer fluorescence lifetime of Cr3+ in the crystal is considered a good candidate for a high powerpulsed laser. However, the phase transition LaAlOs exhi-LaAlO3 compared to Cr3+ in any other laser host material, crystal by many of the popular techniques.

K₂CO₃ concentration used was 7 molal, the pressure was cooling it slowly to 960°C. The flux was poured off and the crystals washed with dilute HNOs. All of the crystals were in the form of rectangular parallelepipeds. Airtron also has produced growth on a seed using a hydrothermal synthesis process where LaAlO3 powder was used for a nutrient. The 20,000 psi, and the growth temperature was approximately a furnace, heated for 16 hr at 1340°C while rotating it and and 81.7 mole % Bi₂O₃ was mixed in proportions with the other oxides so that there were 10.97 and 10.77 moles of $\mathrm{La_2O_3}$ and Al₂O₃ respectively. The platinum crucible was placed in Bi₂O₃-B₂O₃ flux. A Bi₂O₃ flux containing 18.93 mole % B₂O₃ in, have been grown by Airtron(48) in 250-ml crucibles from Single crystals of pale yellow LaAlO₃ measuring $rac{1}{2} imesrac{1}{3} imesrac{1}{4} imesrac{1}{3} imesrac{1}{4}$

crystals up to 43 g in weight were grown and a total of five and Al₂O₃ powder in equal quantities. The melting point of LaAlOs was found to be 2075-2080°C. From this melt, ing from the melt. (49) The crucible was charged with La2O3 Single crystals of LaAlO₃ also have been prepared by pulldifferent boule axis orientations were obtained. 500°C.

GdA103

a temperature of 2030°C while rotating the rod holding the seed at 65 rev/min. Using a pull rate of 6-7 mm/hr crystals $\frac{1}{2}$ in. in diameter and 1 in. long were obtained. The crystals $\frac{1}{2}$ pared gadolinum aluminum oxide, GdAlO₃, single crystals from a melt. The seed was inserted in the melt and pulled at In a study of laser host materials, Mazelsky et al. (50) pretals, however, were not of good optical quality.

 Na_xWO_3

The sodium tungsten bronzes, Na, WO3, were prepared by Straumanis. (31) The method is based on the reaction

 $3x\mathrm{Na_2WO_4} + (6-4x)\mathrm{WO_3} + x\mathrm{W} \longleftarrow 6 \mathrm{Na_xWO_3}.$

To prepare orange to yellow bronzes, perovskite type, in which \hat{x} is 0.6 to 0.9, it was necessary to use mole ratios of reactants 4:2:1 to 9:2:1. The mixture was heated to 850°C

in an inert atmosphere and then slowly cooled. The crystals were recovered by leaching the solid in boiling water, sodium hydroxide solution and then HF

Pb(B233B0.67)Os

ferroelectric compounds Pb(Ni_{0.35}Ta_{0.67})O₃, Pb(Mg_{0.35}Ta_{0.67})O₃ Pb(Co_{0.35}Nb_{0.67})O₃, Pb(Co_{0.35}Nb_{0.67})O₃, Pb(Zn_{0.35}Nb_{0.67})O₃ using a lead oxide flux. A mixture of 60-80 mole % PbO and reagent grade oxides in proper proportions was placed in a platinum crucible and heated to 1200-1300°C. The melt was cooled at a rate of between 30-100°C/hr to a temperature Crystals were separated by boiling in 20% HNO₃ except for The crystals were in the form of imperfect cubes, $1-2~\mathrm{mm}$ $\mathrm{Pb}(\mathrm{Zn_{0.33}Nb_{0.07}})\mathrm{O_3}$ crystals which were washed in acetic acid. of 800°C and then cooled more rapidly to room temperature. Bokov and Myl'nikova(62) prepared single crystals of the

Later, Bokov et al. (33) grew $Pb(Co_{0.5}W_{0.5})O_3$ single crystals using a similar technique. Cobalt carbonate, H_2WO_4 and PbO_4 20-30 mole % WO_3 , and 70-80% PbO. The soaking temperature was 1200°C and the cooling rate to 800°C was were mixed in amounts corresponding to 10 mole % CoO, 5°C/hr. The crystals formed were small cubes. on edge.

Ba(Bo.33 Tao.67)O3

Single crystals of $Ba(B_{0.53}^2Ta_{0.67})O_3$ type compounds⁽⁵⁴⁾ where B^{2+} is Ca, Mg, Zn or Ni were grown by Galasso and Pinto using a BaF_2 flux. The details of the process for each compound and results are presented in Table 10.1.

Pb(B3+B5+5)O3

pared by Galasso and Darby'65) using PbO and PbO-PbF2 fluxes. The conditions of growth and the results are as in Ferroelectric crystals of $Pb(B_0^3;B_0^5;0)O_3$ type compounds, where B^{3+} is Sc or Fe and B^{5+} is Nb or Ta, were pre-Table 10.2.

Table 10.1. Crystal Growth Data for Ba(B_{0.53} Ta_{0.67})O₃-type Com-nounds (after Galasso and Pinto(54))

	Color	yellow yellow green red
	Orystal tal size (mm)	1.0 2.0 0.5 1.5
pounds (after Gaiasso and 1 mio)	Cool- Crysing tal rate size (°C/hr) (mm)	113 100 100
יייייייייייייייייייייייייייייייייייייי	Flux wt(g)	24.0 43.0 18.0 65.0
0000000	Sam- ple wt(g)	6.4 6.4 6.3 11.2
rer or	Max. Soak- Sam- temp. ing ple (°C) time wt(g)	1365 1.0 hr 1400 8.5 hr 1360 0.5 hr 1380 2.0 hr
nas (a	Max. temp. (°C)	1365 1400 1360 1380
nod	Compound	Ba(Ca _{0.38} Ta _{0.67})O ₃ Ba(Mg _{0.38} Ta _{0.67})O ₃ Ba(Ni _{0.38} Ta _{0.67})O ₃ Ba(Zn _{0.38} Ta _{0.67})O ₃

TABLE 10.2. Crystal Growth Data for Pb(B_{0.5}B_{0.5})O₃-type Compounds (after Galasso and Darby⁽⁵⁶⁾)

Crystal size (mm) on edge	
Cooling rate (°C/hr)	30 25
Temp range (°C)	1230-800 1230-800 1160-900 1325-1025
Flux (wt%)	64 54 86 42.5
Flux	PbO PbO PbO PbO- Pbo-
Compound	Pb(Fe, Mb, s)Os Pb(Fe, Ta, s)Os Pb(Sc, sNb, s)Os Pb(Sc, sTa, s)Os

Ba(B3+Ta0.8)0s

num crucible, soaking it for 12 hr at 1470°C and cooling flux. Crystals of Ba(Yo.,Tao.,s)O, up to 0.5 cm on edge were formed by mixing 566.3 g BaCO₃, 164.1 g of Ta₂O₆, 83.9 g $B^{3+} = La$, Gd, Lu, Sc and Y were grown using BaF_2 flux, but more satisfactory results were obtained with B2O3 as the Y₂O₃ and 112 g B₂O₃, placing the mixture in a 250-ml platiit at 1.3°C/hr to 1110°C. A polished crystal is shown in Fig. laser application. (56) Single crystals of these compounds with Galasso et al. studied Ba(B₀.5,Ta_{0.8})O₃-type materials for

rature gradient using a seed crystal suspended below the Growth of $\mathrm{Ba}(\mathrm{Y}_{0.5}\mathrm{Ta}_{0.5})\mathrm{O}_3$ crystals from a flux in a tempePREPARATION OF PEROVSKITE-TYPE OXIDES

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grown) is available in the bottom of the crucible, which is maintained at a higher temperature than the surface of the melt. When the seed crystal is placed in the cooler region of melt surface also was attempted. In this method, an excess of nutrient material (i.e. material of the composition to be the crucible, some of the nutrient dissolves, is transported by diffusion through the flux, and deposits on the rotating seed as it is slowly withdrawn from the melt.

of 0.025 mm/hr. The material which was grown showed large single crystal regions; however, with better control of the volatilization of $\bar{\mathbf{B}}_2\mathbf{O}_3$ large single crystals could probably be 100 rev/min and withdrawn at a rate of 0.0025 in./hr. In a 4-day growth period, the linear growth rate was of the order BaO : $\mathrm{YTaO_4}$: $\mathrm{B_2^{\circ}O_3} = 52.35.13$ was equilibrated for $20\,\mathrm{hr}$. The cible. The seed was lowered into the melt, rotated at about In these experiments a 400-g charge of composition melt surface was held 8°C cooler than the bottom of the cru-

material it purifies the rod as it transforms it into a single through a ring (cold cathode) which impinges electrons on a small zone and melts it. As the molten zone passes along the is similar to an electron beam zone melter except that it requires no anode wire around the insulator and it can be used in partial pressures of oxygen or other gases. The material to be used is made into a polycrystalline rod and passed crystal. The technique seems to be well suited for high meltto grow ${\rm Al_2O_3}$ single crystals. The apparatus (see Fig. 10.10)Another technique(66) considered was one developed with J. Davis, also of the United Aircraft Research Laboratories ing point oxides. grown.

La(B_{0.5} B_{0.5} O₃

B2+, is Ni, Mg, Zn and B4+ is Ir or Ru were grown from fluxes in platinum crucibles (55). The amount and composition Single crystals of La(B2+B4+)Os-type compounds where of the fluxes, firing conditions and cooling rates are listed in Table 10.3.

The crystals produced in this manner were found to be electrically conducting.

Table 10.3. Crystal Growth Data for La(B2+B4+)O3-type Compounds (after Galasso and Darby(55))

Compound	Flux	Flux (wt%)	Temp. range (°C)	Cooling rate (°C/hr)	Crystal size (mm) on edge
La(Mg., Ruo., s)Os	Pb0-	82	1320-1000	30	1.0
La(Nio.sIro.s)Os	Pbr. Pbo-	85	1300-1000	30	0.5
La(Ni _{0.8} Ru _{0.8})O ₃	-Pb0-	86	1300-1000	30	2.0
La(Zn _{0.6} Ru _{0.6})O ₃	Pbr.	8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8	1300-850 1300-25	30	0.0

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CHAPTER 11

OTHER PEROVSKITE-TYPE COMPOUNDS

The oxides are by far the most numerous and most interesting materials with the perovskite structure. However, there are some carbides, halides, hydrides and nitrides with this structure which have been investigated because of their magnetic properties and possible use as hosts for transition metal activating ions. Some of the data on their preparation, structure and properties is presented in this chapter.

11.1. Preparation of Perovskite-type Phases (other than oxides)

The ternary carbides with the perovskite structure have been prepared predominantly by two techniques. The first involves melting the appropriate proportions of the two metals and carbon under argon and cooling them. Whenever pronounced coring exists, the samples are annealed for long periods. (1) The second technique is a solid-state reaction between the reactants after they are placed in an evacuated sealed silica capsule. (2) In a variation of this method, one metal is heated with carbon and the alloy is ground, mixed with the other metal and heated again.

Perovskite-type fluorides have been prepared by precipitation from aqueous solutions. However, crystals prepared pitation from aqueous solutions. However, crystals prepared in this manner are not stoichiometric. (3) Therefore, other techniques are used when possible. The halides K(Na_{0.5}Gr_{0.5})F₃, niques are used when possible. The halides K(Na_{0.5}Gr_{0.5})F₃ were formed by react-K(Na_{0.5}Fe_{0.5})F₃ and K(Na_{0.5}Ga_{0.5})F₂ were formed by reacting the trifluorides with KHF₃ and NaHF₂ in a platinum crucible over an open flame. (4) The melt was cooled, leached with water and ethyl alcohol, and the product dried.

Many fluorides with the perovskite structure also have been prepared by a solid-state reaction between an alkali metal halide and a divalent metal oxide at $500-800^{\circ}\text{C}^{(5)}$. The same type of reactions have been conducted between the binary fluorides.

Single crystals of a number of fluorides have been prepared by Knox using fluorides precipitated from aqueous solutions as the reactants. (a) The fluorides were heated in anhydrous HF, mixed with KHF2, melted and slowly cooled in an inert atmosphere. Crystals of KMnF3, KFeF3, KCoF3, KNiF3, KCuF3 and KZnF3 have been grown in this manner.

Single crystals of KMnF₃ have been grown by the Czochralski technique by Nassau. (**) In these studies, special equip-

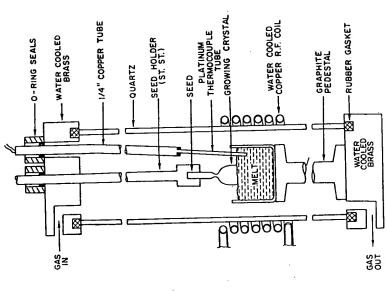


Fig. 11.1. Apparatus for Czochralski-pulling (after Nassau $^{(n)}$).

ment had to be fabricated to prevent any oxygen from reacting with the melt (see Fig. 11.1)

were used in dense graphite containers under an HF-argon atmosphere. Single crystals 2×0.5 in. in dimensions were Kestigian et al. (3) grew crystals of RbFeF, and CsFeF, using a horizontal Bridgman technique. Anhydrous fluorides

gen atmosphere.(9) The metals were ground and treated in a by placing mixtures of the metals in a stainless-steel boat in The ternary hydrides, LiBaH, and LiSrH, were prepared a stainless-steel bomb and heating the mixture under a hydrogrown by this method.

ground into powders and a mixture of NH3 and hydrogen that they could obtain nitrides by first processing the ingots was used as the nitriding gas. Weiner and Berger(11) found into strips before nitriding. If the strips were very thin, hopared by Stadelmaier and Fraker(10) using an induction unit to melt the alloy, which they then nitrided. The alloy was The nitrides Fe₃NiN, Fe₃PtN, Fe₄N and Fe₃PdN were predry box under an argon atmosphere. mogeneous nitrides were obtained.

11.2. STRUCTURE

produced by a face centered cubic arrangement of atoms with a superstructure. In the unit cell of Mn₃AlC structure the manganese atoms are located at the face centers, the aluminum atom is at the cube corners and the carbon atom is In the structure of the ternary carbides described in this the C atom in the B position, and the transition metal atoms This makes the X-ray pattern of these phases look like that in the oxygen atom positions of the perovskite structure. chapter the Al, Sn or Ga metal atoms are in the A position,

cause of the low scattering factor of nitrogen the powder patterns of these compounds also look as though they had a The nitrides Fe₃NiN and Fe₃PtN also adopt the perovskite structure. Studies by Wiener and Berger indicate that there is complete ordering in the structure of these nitrides. Bestructure with a face-centered cubic lattice. at the body centered position.

Table 11.1. Unit Cell Parameters for Perovskite-type Phases

OTHER PEROVSKITE-TYPE COMPOUNDS

(other than oxides)

Refer- ence s	12 1 2 2 2	13 13 13, 14 13, 14	, , , , , , , , , , , , , , , , , , ,
		hexagonal tetragonal tetragonal	tetragonal tetragonal tetragonal tetragonal
c (A)		14.866 8.72 8.39	5.630 9.05 4.019 3.926 8.76
a (Å)	3.719 3.869 3.86 8.376 3.92	4.522 10.70 5.20 6.158 5.47 5.77 5.44	5.874 5.590 9.90 8.742 4.293 4.140 4.120 3.996 7.76 4.062 4.062 4.062 8.19
Phases	Carbides AlFe ₃ C AlMn ₃ C Fe ₃ SnC GaMn ₃ C Mn ₃ ZnC	Halides CsCaBr _s CsCdBr _s CsCdCl _s CsFeF _s CsGeCl _s CaBgBr _s	CSPDBT; CSPDBT; CSPDBT; CSZDF; KCGF; KCGF; KCOF; KCOF; KKNF; KKNF; KNIF; KNIF; KNIF; KNSF; KNOSP; KNOSP; KNOSP; KNOSP; KNOSP; KNOSP; KNOSP; KNOSP; KNOSP; KNOSP; KNOSP; KNOSP; KNOSP;

Refer- ences	L 2 4 4 4	9 9 9	1111
	tetragonal		
c (Å)	8.01		
d (A)	4.250 8.71 8.266 8.323 8.246	4.023 3.796 3.833	3.795 3.867 3.790 3.857
Phases	RbMnF ₃ RbZnF ₈ K(Cr ₀ ·s/Na ₀ ·s/F ₃ K(Fe ₀ ·s/Na ₀ ·s/F ₃ K(Ga ₀ ·s/Na ₀ ·s/F ₃	Hydrides LiBaH, LiEuH, LiSrH,	Nitrides Fe ₄ N Mn ₄ N Fe ₃ NiN Fe ₃ PtN

As can be seen in Table 11.1, many of the fluorides have the "ideal" cubic perovskite structure. Of the series with the KCuF, and KCrF, have distorted structures. Many of the formula KBF₃(B²⁺ = Mn, Fe, Co, Ni, Cu, Cr and Zn), only others have structures related to the various modification of the perovskite type.

were found to adopt the ordered perovskite structure of the (NH4)3FeF6 type. The fluoride ions in this structure move The complex fluorides $K(Na_{0.5}Cr_{0.5})F_3$ and $K(Na_{0.5}Fe_{0.5})\overline{F}_3$ closer to the transition metal ion and away from the sodium

perovskite structure with the lithium ions in the B position The ternary hydrides LiBaH₃ and LiSrH₃ have the inverse and alkaline earth metal ions in the A position. The crystallographic data for these phases as well as for the carbides. halides and nitrides are listed in Table 11.1.

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was ferromagnetic with a Curie temperature of 80°C. The Curie temperatures of both of these materials varied with the Mn/Al or Zn ratio. Figure 11.2 shows the variation of Curie structure are ferromagnetic materials. The carbide Mn₃AlC has been studied extensively. Butters and Myers⁽²⁾ found that it was strongly magnetic at low temperature and has a Curie temperature of 15°C. They also found that Mn₃ZnC Many of the carbides and nitrides with the perovskite

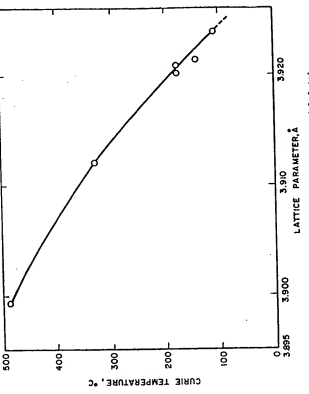


Fig. 11.2. Variation of Curie temperature with lattice parameters (after Butters and Myers⁽²⁾)

temperature with lattice constant for the Mn,Zn,C phases. Note that the Curie temperature increases as the lattice constant decreases.

at 20° and decreases with increasing temperaul.0 It is not The electrical resistivity of Mn_3ZnC is 770 $\times cat^6$ ohm-cm stable in moist air and must be kept in a desictor.re

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The nitrides Fe,N, Mn,N, Fe,NiN and Fe,PtN were also found to be ferromagnetic with Curie points of 488°, 465°, 487° and 369°C, respectively

215°K for KCuF, appears to be less certain. However, a study of the absorption spectra of KMnF₃ as a function of ferromagnetic with Néel points of 112°, 135° and 280°K. The evidence for antiferromagnetism below 80°K for KMnF3 and temperature does show anomalies in the maxima at $184^\circ
m K$ nary perovskite-type fluorides also had interesting magnetic properties. The fluorides KFeF3, KCoF3 and KNiF3 are antiand at 88°K. The first the authors attribute to a phase trans-Studies by Machin et al. (19) showed that many of the terformation and the latter to electron spin coupling.

75° and 83°K respectively. Small changes in the fluorescent reported for the latter compound and KMnF3 (22, 23) At about half the Néel temperatures larger and more strongly cence of these materials. This effect may involve the coupling Other fluorides which were found to be antiferromagnetic are RbFeF₃⁽²⁰⁾ and RbMnF₃, (21) with Néel temperatures of properties in the vicinity of the Néel temperature have been temperature-dependent changes were observed in the fluoresof the lattice and magnetic interaction of the excited Mn^{2+}

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crystallography

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